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(54) Title: FLUORESCENT PROTEINS FROM NON-BIOLUMINESCENT SPECIES OF CLASS ANTHOZOA, GENES ENCODING SUCH PROTEINS AND USES THEREOF		
(57) Abstract		
<p>The present invention is directed to novel fluorescent proteins from non-bioluminescent organisms from the Class Anthozoa. Also disclosed are methods of identifying nucleic acid sequence encoding the fluorescent proteins and further analyzing the proteins.</p>		

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**FLUORESCENT PROTEINS FROM NON-BIOLUMINESCENT SPECIES
OF CLASS ANTHOZOA, GENES ENCODING SUCH PROTEINS AND
USES THEREOF**

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BACKGROUND OF THE INVENTION

Field of the Invention

This invention relates to the field of molecular biology. More specifically, this invention relates to novel fluorescent proteins, methods of identifying the DNA sequences encoding the proteins and uses thereof.

Description of the Related Art

Fluorescence labeling is a particularly useful tool for marking a protein, cell, or organism of interest. Traditionally, a protein of interest is purified, then covalently conjugated to a fluorophore derivative. For *in vivo* studies, the protein-dye complex is then inserted into cells of interest using micropipetting or a method of reversible permeabilization. The dye attachment and insertion steps, however, make the process laborious and difficult to control. An alternative method of labeling proteins of interest is to concatenate or fuse the gene expressing the protein of interest to a gene expressing a marker, then express the fusion product. Typical markers for this method of protein labeling include β -galactosidase, firefly luciferase

and bacterial luciferase. These markers, however, require exogenous substrates or cofactors and are therefore of limited use for *in vivo* studies.

A marker that does not require an exogenous cofactor or 5 substrate is the green fluorescent protein (GFP) of the jellyfish *Aequorea victoria*, a protein with an excitation maximum at 395 nm, a second excitation peak at 475 nm and an emission maximum at 510 nm. GFP is a 238-amino acid protein, with amino acids 65-67 involved in the formation of the chromophore.

10 Uses of GFP for the study of gene expression and protein localization are discussed in detail by Chalfie et al. in *Science* 263 (1994), 802-805, and Heim et al. in *Proc. Nat. Acad. Sci.* 91 (1994), 12501-12504. Additionally, Rizzuto et al. in *Curr. Biology* 5 (1995), 635-642, discuss the use of wild-type GFP as a tool for visualizing 15 subcellular organelles in cells, while Kaether and Gerdes in *Febs Letters* 369 (1995), 267-271, report the visualization of protein transport along the secretory pathway using wild-type GFP. The expression of GFP in plant cells is discussed by Hu and Cheng in *Febs Letters* 369 (1995), 331-334, while GFP expression in *Drosophila* embryos is described by 20 Davis et al. in *Dev. Biology* 170 (1995), 726-729.

Crystallographic structures of wild-type GFP and the mutant GFP S65T reveal that the GFP tertiary structure resembles a barrel 25 (Ormö et al., *Science* 273 (1996), 1392-1395; Yang, et al., *Nature Biotechnol* 14 (1996), 1246-1251). The barrel consists of beta sheets in a compact structure, where, in the center, an alpha helix containing the chromophore is shielded by the barrel. The compact structure makes GFP very stable under diverse and/or harsh conditions such as protease treatment, making GFP an extremely useful reporter in

general. However, the stability of GFP makes it sub-optimal for determining short-term or repetitive events.

A great deal of research is being performed to improve the properties of GFP and to produce GFP reagents useful and optimized for 5 a variety of research purposes. New versions of GFP have been developed, such as a "humanized" GFP DNA, the protein product of which has increased synthesis in mammalian cells (Haas, et al., *Current Biology* 6 (1996), 315-324; Yang, et al., *Nucleic Acids Research* 24 (1996), 4592-4593). One such humanized protein is "enhanced green 10 fluorescent protein" (EGFP). Other mutations to GFP have resulted in blue-, cyan- and yellow-green light emitting versions. Despite the great utility of GFP, however, other fluorescent proteins with properties similar to or different from GFP would be useful in the art. Novel 15 fluorescent proteins result in possible new colors, or produce pH-dependent fluorescence. Other benefits of novel fluorescent proteins include fluorescence resonance energy transfer (FRET) possibilities based on new spectra and better suitability for larger excitation.

The prior art is deficient in novel fluorescent proteins wherein the DNA coding sequences are known. The present invention 20 fulfills this long-standing need in the art.

SUMMARY OF THE INVENTION

25 The present invention is directed to an isolated and purified fluorescent protein selected from the group consisting of amFP486, cFP484, zFP506, zFP538, dsFP483, drFP583, asFP600, dgFP512 and dmFP592.

In one embodiment of the present invention, there is provided a method of identifying a DNA sequence encoding a fluorescent protein comprising the step of screening for an existence of a nucleic acid sequence in a sample, wherein the nucleic acid sequence encodes a peptide having a sequence selected from the group consisting of SEQ ID Nos. 3, 5, 8, 11, 12 and 14. The existence of the nucleic acid sequence identifies the DNA sequence encoding the fluorescent protein.

In another embodiment of the present invention, there is provided a method of identifying a DNA sequence encoding a fluorescent protein comprising the step of screening for an existence of a nucleic acid sequence in a sample, wherein the nucleic acid sequence hybridizes to a primer selected from the group consisting of SEQ ID Nos. 4, 6, 7, 9, 10, 13, 15 and 16. The existence of the nucleic acid sequence identifies the DNA sequence encoding the fluorescent protein.

In still another embodiment of the present invention, there is provided a method of analyzing a fluorescent protein in a cell, comprising the steps of expressing a nucleic acid sequence encoding a fluorescent protein having an amino acid sequence selected from the group consisting of SEQ ID Nos. 55-63 in the cell; and measuring a fluorescence signal from the protein. This method further comprises a step of sorting the cell according to the signal. Preferably, the cell is sorted by fluorescence activated cell sorting. Still preferably, the nucleic acid sequence comprises a gene of interest encoding a protein of interest fused to the fluorescent protein, wherein the protein of interest is distinct from the fluorescent protein. The detected fluorescence signal indicates the presence of the gene of interest and further the protein of interest in the cell. By identifying an

intracellular location of the fluorescent protein, an intracellular location of the protein of interest is also identified.

Other and further aspects, features, and advantages of the present invention will be apparent from the following description of 5 the presently preferred embodiments of the invention given for the purpose of disclosure.

BRIEF DESCRIPTION OF THE DRAWINGS

10 **Figure 1** shows the modified strategy of 3'-RACE used to isolate the target fragments. Sequences of the oligonucleotides used are shown in Table 2. Dp1 and Dp2 are the degenerate primers used in the first and second PCR, respectively (see Tables 3 and 4 for the sequences of degenerate primers).

15 **Figure 2A** shows multiple alignment of novel fluorescent proteins. The numbering is based on *Aequorea victoria* green fluorescent protein (GFP). Two proteins from *Zoanthus* and four from *Discosoma* are compared between each other: residues identical to the corresponding ones in the first protein of the series are represented by 20 dashes. Introduced gaps are represented by dots. In the sequence of *A. victoria* GFP, the stretches forming beta-sheets are underlined; the residues whose side chains form the interior of the beta-can are shaded (according to Yang et al., *Nature Biotechnol.* 14, 1246–1251 (1996)).
25 **Figure 2B** shows the N-terminal part of cFP484, which has no homology with the other proteins. The putative signal peptide is underlined.

Figure 3 shows the excitation and emission spectrum of the novel fluorescent protein from *Anemonia majano*, amFP486.

Figure 4 shows the excitation and emission spectrum of the novel fluorescent protein from *Clavularia*, cFP484.

Figure 5 shows the excitation and emission spectrum of the novel fluorescent protein from *Zoanthus*, zFP506.

5 **Figure 6** shows the excitation and emission spectrum of the novel fluorescent protein from *Zoanthus*, zFP538.

Figure 7 shows the excitation and emission spectrum of the novel fluorescent protein from *Discosoma striata*, dsFP483.

10 **Figure 8** shows the excitation and emission spectrum of the novel fluorescent protein from *Discosoma*, drFP583.

Figure 9 shows the excitation and emission spectrum of the novel fluorescent protein from *Anemonia sulcata*, asFP600.

Figure 10 shows the excitation and emission spectrum of the novel fluorescent protein from *Discosoma*, dgFP512.

15 **Figure 11** shows the excitation and emission spectrum of the novel fluorescent protein from *Discosoma*, dmFP592.

DETAILED DESCRIPTION OF THE INVENTION

20 As used herein, the term "GFP" refers to the basic green fluorescent protein from *Aequorea victoria*, including prior art versions of GFP engineered to provide greater fluorescence or fluoresce in different colors. The sequence of *Aequorea victoria* GFP (SEQ ID No. 54) has been disclosed in Prasher et al., *Gene* 111 (1992), 229-33.

25 As used herein, the term "EGFP" refers to mutant variant of GFP having two amino acid substitutions: F64L and S65T (Heim et al., *Nature* 373 (1995), 663-664). The term "humanized" refers to changes made to the GFP nucleic acid sequence to optimize the codons for

expression of the protein in human cells (Yang et al., *Nucleic Acids Research* 24 (1996), 4592-4593).

In accordance with the present invention there may be employed conventional molecular biology, microbiology, and recombinant DNA techniques within the skill of the art. Such techniques are explained fully in the literature. See, e.g., Maniatis, Fritsch & Sambrook, "Molecular Cloning: A Laboratory Manual" (1982); "DNA Cloning: A Practical Approach," Volumes I and II (D.N. Glover ed. 1985); "Oligonucleotide Synthesis" (M.J. Gait ed. 1984); "Nucleic Acid Hybridization" (B.D. Hames & S.J. Higgins eds. (1985)); "Transcription and Translation" (B.D. Hames & S.J. Higgins eds. (1984)); "Animal Cell Culture" (R.I. Freshney, ed. (1986)); "Immobilized Cells and Enzymes" (IRL Press, (1986)); B. Perbal, "A Practical Guide To Molecular Cloning" (1984).

A "vector" is a replicon, such as plasmid, phage or cosmid, to which another DNA segment may be attached so as to bring about the replication of the attached segment.

A "DNA molecule" refers to the polymeric form of deoxyribonucleotides (adenine, guanine, thymine, or cytosine) in either single stranded form or a double-stranded helix. This term refers only to the primary and secondary structure of the molecule, and does not limit it to any particular tertiary forms. Thus, this term includes double-stranded DNA found, inter alia, in linear DNA molecules (e.g., restriction fragments), viruses, plasmids, and chromosomes.

A DNA "coding sequence" is a DNA sequence which is transcribed and translated into a polypeptide *in vivo* when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a start codon at the 5' (amino) terminus and a translation stop codon at the 3'

(carboxyl) terminus. A coding sequence can include, but is not limited to, prokaryotic sequences, cDNA from eukaryotic mRNA, genomic DNA sequences from eukaryotic (e.g., mammalian) DNA, and synthetic DNA sequences. A polyadenylation signal and transcription termination 5 sequence may be located 3' to the coding sequence.

As used herein, the term "hybridization" refers to the process of association of two nucleic acid strands to form an antiparallel duplex stabilized by means of hydrogen bonding between residues of the opposite nucleic acid strands.

10 The term "oligonucleotide" refers to a short (under 100 bases in length) nucleic acid molecule.

"DNA regulatory sequences", as used herein, are transcriptional and translational control sequences, such as promoters, enhancers, polyadenylation signals, terminators, and the like, that 15 provide for and/or regulate expression of a coding sequence in a host cell.

A "promoter sequence" is a DNA regulatory region capable of binding RNA polymerase in a cell and initiating transcription of a downstream (3' direction) coding sequence. For purposes of defining 20 the present invention, the promoter sequence is bounded at its 3' terminus by the transcription initiation site and extends upstream (5' direction) to include the minimum number of bases or elements necessary to initiate transcription at levels detectable above background. Within the promoter sequence will be found a 25 transcription initiation site, as well as protein binding domains responsible for the binding of RNA polymerase. Eukaryotic promoters will often, but not always, contain "TATA" boxes and "CAT" boxes. Various promoters, including inducible promoters, may be used to drive the various vectors of the present invention.

As used herein, the terms "restriction endonucleases" and "restriction enzymes" refer to bacterial enzymes, each of which cut double-stranded DNA at or near a specific nucleotide sequence.

A cell has been "transformed" or "transfected" by 5 exogenous or heterologous DNA when such DNA has been introduced inside the cell. The transforming DNA may or may not be integrated (covalently linked) into the genome of the cell. In prokaryotes, yeast, and mammalian cells for example, the transforming DNA may be maintained on an episomal element such as a plasmid. With respect to 10 eukaryotic cells, a stably transformed cell is one in which the transforming DNA has become integrated into a chromosome so that it is inherited by daughter cells through chromosome replication. This stability is demonstrated by the ability of the eukaryotic cell to establish cell lines or clones comprised of a population of daughter 15 cells containing the transforming DNA. A "clone" is a population of cells derived from a single cell or common ancestor by mitosis. A "cell line" is a clone of a primary cell that is capable of stable growth *in vitro* for many generations.

A "heterologous" region of the DNA construct is an 20 identifiable segment of DNA within a larger DNA molecule that is not found in association with the larger molecule in nature. Thus, when the heterologous region encodes a mammalian gene, the gene will usually be flanked by DNA that does not flank the mammalian genomic DNA in the genome of the source organism. In another example, 25 heterologous DNA includes coding sequence in a construct where portions of genes from two different sources have been brought together so as to produce a fusion protein product. Allelic variations or naturally-occurring mutational events do not give rise to a heterologous region of DNA as defined herein.

As used herein, the term "reporter gene" refers to a coding sequence attached to heterologous promoter or enhancer elements and whose product may be assayed easily and quantifiably when the construct is introduced into tissues or cells.

5 The amino acids described herein are preferred to be in the "L" isomeric form. The amino acid sequences are given in one-letter code (A: alanine; C: cysteine; D: aspartic acid; E: glutamic acid; F: phenylalanine; G: glycine; H: histidine; I: isoleucine; K: lysine; L: leucine; M: methionine; N: asparagine; P: proline; Q: glutamine; R: arginine; S: serine; T: threonine; V: valine; W: tryptophane; Y: tyrosine; X: any residue). NH₂ refers to the free amino group present at the amino terminus of a polypeptide. COOH refers to the free carboxy group present at the carboxy terminus of a polypeptide. In keeping with standard polypeptide nomenclature, *J Biol. Chem.*, 243 (1969), 3552-10 59 is used.
15

The present invention is directed to an isolated and purified fluorescent protein selected from the group consisting of amFP486, cFP484, zFP506, zFP538, dsFP483, drFP583, asFP600, dgFP512 and dmFP592.

20 In one embodiment of the present invention, there is provided a method of identifying a DNA sequence encoding a fluorescent protein comprising the step of screening for an existence of a nucleic acid sequence in a sample, wherein the nucleic acid sequence encodes a peptide having a sequence selected from the group 25 consisting of SEQ ID Nos. 3, 5, 8, 11, 12 and 14. The existence of the nucleic acid sequence identifies the DNA sequence encoding the fluorescent protein.

In another embodiment of the present invention, there is provided a method of identifying a DNA sequence encoding a

fluorescent protein comprising the step of screening for an existence of a nucleic acid sequence in a sample, wherein the nucleic acid sequence hybridizes to a primer selected from the group consisting of SEQ ID Nos. 4, 6, 7, 9, 10, 13, 15 and 16. The existence of the nucleic acid 5 sequence identifies the DNA sequence encoding the fluorescent protein.

In still another embodiment of the present invention, there is provided a method of analyzing a fluorescent protein in a cell, comprising the steps of expressing a nucleic acid sequence encoding a 10 fluorescent protein having an amino acid sequence selected from the group consisting of SEQ ID Nos. 55-63 in the cell; and measuring a fluorescence signal from the protein. This method further comprises a step of sorting the cell according to the signal. Preferably, the cell is sorted by fluorescence activated cell sorting. Still preferably, the 15 nucleic acid sequence comprises a gene of interest encoding a protein of interest fused to the fluorescent protein, wherein the protein of interest is distinct from the fluorescent protein. The detected fluorescence signal indicates the presence of the gene of interest and further the protein of interest in the cell. By identifying an 20 intracellular location of the fluorescent protein, an intracellular location of the protein of interest is also identified.

The following examples are given for the purpose of illustrating various embodiments of the invention and are not meant to limit the present invention in any fashion.

EXAMPLE 1**Biological Material**

5 Novel fluorescent proteins were identified from several genera of Anthozoa which do not exhibit any bioluminescence but have fluorescent color as observed under usual white light or ultraviolet light. Six species were chosen (see Table 1).

10

TABLE 1**Anthozoa Species Used in This Study**

Species	Area of Origination	Fluorescent Color
Anemonia majano	Western Pacific	bright green tentacle tips
Clavularia sp.	Western Pacific	bright green tentacles and oral disk
Zoanthus sp.	Western Pacific	green-yellow tentacles and oral disk
Discosoma sp. "red"	Western Pacific	orange-red spots oral disk
Discosoma sp. striata	Western Pacific	blue-green stripes on oral disk
Discosoma sp. "magenta"	Western Pacific	faintly purple oral disk
Discosoma sp.	Western Pacific	green spots on oral disk

“green”		
Anemonia sulcata	Mediterranean	purple tentacle tips

EXAMPLE 2**cDNA Preparation**

Total RNA was isolated from the species of interest according to the protocol of Chomczynski and Sacchi (Chomczynski P., et al., *Anal. Biochem.* 162 (1987), 156-159). First-strand cDNA was synthesized starting with 1-3 µg of total RNA using SMART PCR cDNA synthesis kit (CLONTECH) according to the provided protocol with the only alteration being that the "cDNA synthesis primer" provided in the kit was replaced by the primer TN3 (5'- CGCAGTCGACCG(T)₁₃, SEQ ID No. 1) (Table 2). Amplified cDNA samples were then prepared as described in the protocol provided except the two primers used for PCR were the TS primer (5'-AAGCAGTGGTATCAACGCAGAGT, SEQ ID No. 2) (Table 2) and the TN3 primer (Table 2), both in 0.1 µM concentration. Twenty to twenty-five PCR cycles were performed to amplify a cDNA sample. The amplified cDNA was diluted 20-fold in water and 1 µl of this dilution was used in subsequent procedures.

TABLE 2Oligos Used in cDNA Synthesis and RACE

5 TN3: 5'-CGCAGTCGACCG(T)₁₃
(SEQ ID No. 1)

T7-TN3: 5'-GTAATACGACTCACTATAAGGCCGCAGTCGACCG(T)₁₃
(SEQ ID No. 17)

10 TS-primer: 5'-AAGCAGTGGTATCAACGCAGAGT
(SEQ ID No. 2)

15 T7-TS:
5'-GTAATACGACTCACTATAAGGCAGCAGTGGTATCAACGCAGAGT
(SEQ ID No. 18)

T7: 5'-GTAATACGACTCACTATAAGGC
(SEQ ID No. 19)

20 TS-oligo 5'-AAGCAGTGGTATCAACGCAGAGTACGCrGrGrG
(SEQ ID No. 53)

25

EXAMPLE 3**Oligo Design**

To isolate fragments of novel fluorescent protein cDNAs,
5 PCR using degenerate primers was performed. Degenerate primers
were designed to match the sequence of the mRNAs in regions that
were predicted to be the most invariant in the family of fluorescent
proteins. Four such stretches were chosen (Table 3) and variants of
10 degenerate primers were designed. All such primers were directed to
the 3'-end of mRNA. All oligos were gel-purified before use. Table 2
shows the oligos used in cDNA synthesis and RACE.

TABLE 3

Key Amino Acid Stretches and Corresponding Degenerate Primers Used for Isolation of Fluorescent Proteins

5

Stretch Position according to A. victoria GFP (7)	Amino Acid Sequence of the Key Stretch	Degenerated Primer Name and Sequence
20-25	GXVN ^G H (SEQ ID No. 3)	NGH: 5'- GA(C,T) GGC TGC GT(A,T,G,C) AA(T,C) GG(A,T,G) CA (SEQ ID No. 4)
31-35	GEGEG (SEQ ID No. 5) GEGNG (SEQ ID No. 8)	GEGA: 5'- GTT ACA GGT GA(A,G) GG(A,C) GA(A,G) GG (SEQ ID No. 6) GEGb: 5'- GTT ACA GGT GA(A,G) GG(T,G) GA(A,G) GG (SEQ ID No. 7) GNGa: 5'- GTT ACA GGT GA(A,G) GG(A,C) AA(C,T) GG (SEQ ID No. 9) GNGb: 5'- GTT ACA GGT GA(A,G) GG(T,G) AA(C,T) GG (SEQ ID No. 10)
127-131	GMNFP (SEQ ID No. 11) GVNFP (SEQ ID No. 12)	NFP: 5' TTC CA(C,T) GGT (G,A)TG AA(C,T) TT(C,T) CC (SEQ ID NO. 13)
134-137	GPVM (SEQ ID No. 14)	PVMA: 5' CCT GCC (G,A)A(C,T) GGT CC(A,T,G,C) GT(A,C) ATG (SEQ ID NO. 15) PVMB: 5' CCT GCC (G,A)A(C,T) GGT CC(A,T,G,C) GT(G,T) ATG (SEQ ID NO. 16)

EXAMPLE 4Isolation of 3'-cDNA Fragments of nFPs

The modified strategy of 3'-RACE was used to isolate the target fragments (see Figure 1). The RACE strategy involved two consecutive PCR steps. The first PCR step involved a first degenerate primer (Table 4) and the T7-TN3 primer (SEQ ID No. 17) which has a 3' portion identical to the TN3 primer used for cDNA synthesis (for sequence of T7-TN3, Table 2). The reason for substituting the longer T7-TN3 primer in this PCR step was that background amplification which occurred when using the shorter TN3 primer was suppressed effectively, particularly when the T7-TN3 primer was used at a low concentration (0.1 μ M) (Frohman et al., (1998) *PNAS USA*, 85, 8998-9002). The second PCR step involved the TN3 primer (SEQ ID No. 1, Table 2) and a second degenerate primer (Table 4).

TABLE 4

Combinations of Degenerate Primers for First and Second PCR Resulting in Specific Amplification of 3'-Fragments of nFP cDNA

Species	First Degenerate Primer	Second Degenerate Primer
<i>Anemonia majano</i>	NGH (SEQ ID No. 4)	GNGb (SEQ ID No. 10)
<i>Clavularia</i> sp.	NGH (SEQ ID No. 4)	GEGA (SEQ ID No. 6)
<i>Zoanthus</i> sp.	NGH (SEQ ID No. 4)	GEGA (SEQ ID No. 6)
<i>Discosoma</i> sp. "red"	NGH (SEQ ID No. 4)	GEGA (SEQ ID No. 6), NFP (SEQ ID No. 13) or PVMB (SEQ ID No. 16)
<i>Discosoma striata</i>	NGH (SEQ ID No. 4)	NFP (SEQ ID No. 13)
<i>Anemonia sulcata</i>	NGH (SEQ ID No. 4)	GEGA (SEQ ID No. 6) or NFP (SEQ ID No. 13)

5

The first PCR reaction was performed as follows: 1 μ l of 20-fold dilution of the amplified cDNA sample was added into the reaction mixture containing 1X Advantage KlenTaq Polymerase Mix with provided buffer (CLONTECH), 200 μ M dNTPs, 0.3 μ M of first degenerate

primer (Table 4) and 0.1 μ M of T7-TN3 (SEQ ID No. 17) primer in a total volume of 20 μ l. The cycling profile was (Hybaid OmniGene Thermocycler, tube control mode): 1 cycle for 95°C, 10 sec.; 55°C, 1 min.; 72°C, 40 sec; 24 cycles for 95°C, 10 sec.; 62°C, 30 sec.; 72°C, 40 sec. The reaction was then diluted 20-fold in water and 1 μ l of this dilution was added to a second PCR reaction, which contained 1X Advantage KlenTaq Polymerase Mix with the buffer provided by the manufacturer (CLONTECH), 200 μ M dNTPs, 0.3 μ M of the second degenerate primer (Table 4) and 0.1 μ M of TN3 primer. The cycling profile was (Hybaid OmniGene Thermocycler, tube control mode): 1 cycle for 95°C, 10 sec.; 55°C (for GEG/GNG or PVM) or 52°C (for NFP), 1 min.; 72°C, 40 sec; 13 cycles for 95°C, 10sec.; 62°C (for GEG/GNG or PVM) or 58°C (for NFP), 30 sec.; 72°C, 40 sec. The product of PCR was cloned into PCR-Script vector (Stratagene) according to the manufacturer's protocol.

Different combinations of degenerate primers were tried in the first and second PCR reactions on the DNA from each species until a combination of primers was found that resulted in specific amplification--meaning that a pronounced band of expected size (about 650-800 bp for NGH and GEG/GNG and 350-500 bp for NFP and PVM--sometimes accompanied by a few minor bands) was detected on agarose gel after two PCR reactions. The primer combinations of choice for different species of the Class Anthozoa are listed in Table 4. Some other primer combinations also resulted in amplification of fragments of correct size, but the sequence of these fragments showed no homology to the other fluorescent proteins identified or to *Aequorea victoria* GFP.

EXAMPLE 5

Obtaining Full-Length cDNA Copies

Upon sequencing the obtained 3'-fragments of novel fluorescent protein cDNAs, two nested 5'-directed primers were synthesized for cDNA (Table 5), and the 5' ends of the cDNAs were then amplified using two consecutive PCRs. In the next PCR reaction, the novel approach of "step-out PCR" was used to suppress background amplification. The step-out reaction mixture contained 1x Advantage KlenTaq Polymerase Mix using buffer provided by the manufacturer (CLONTECH), 200 µM dNTPs, 0.2 µM of the first gene-specific primer (see Table 5), 0.02 µM of the T7-TS primer (SEQ ID No. 18), 0.1 µM of T7 primer (SEQ ID No. 19) and 1 µl of the 20-fold dilution of the amplified cDNA sample in a total volume of 20 µl. The cycling profile was (Hybaid OmniGene Thermocycler, tube control mode): 23-27 cycles for 95°C, 10 sec.; 60°C, 30 sec.; 72°C, 40 sec. The product of amplification was diluted 50-fold in water and one µl of this dilution was added to the second (nested) PCR. The reaction contained 1X Advantage KlenTaq Polymerase Mix with provided buffer (CLONTECH), 200 µM dNTPs, 0.2 µM of the second gene-specific primer and 0.1 µM of TS primer (SEQ ID No. 2) in a total volume of 20 µl. The cycling profile was (Hybaid OmniGene Thermocycler, tube control mode): 12 cycles for 95°C, 10 sec.; 60°C, 30 sec.; 72°C, 40 sec. The product of amplification was then cloned into pAtlas vector (CLONTECH) according to the manufacturer's protocol.

TABLE 5Gene-Specific Primers Used for 5'-RACE

Species	First Primer	Second (Nested) Primer
Anemonia majano	5'-GAAATAGTCAGGCATACTGGT (SEQ ID No. 20)	5'-GTCAGGCATAC TGGTAGGAT (SEQ ID No. 21)
Clavularia sp.	5'-CTTGAAATAGTCTGCTATATC (SEQ ID No. 22)	5'-TCTGCTATATC GTCTGGGT (SEQ ID No. 23)
Zoanthus sp.	5'- GTTCTGAAATAGTCTACTATGT (SEQ ID No. 24)	5'-GTCTACTATGTCTT GAGGAT (SEQ ID No. 25)
Discosoma sp. "red"	5'-CAAGCAAATGGCAAAGGTC (SEQ ID No. 26)	5'-CGGTATTGTGGCC TTCGTA (SEQ ID No. 27)
Discosoma striata	5'-TTGTCTTCTTCTGCACAAC (SEQ ID No. 28)	5'-CTGCACAACGG GTCCAT (SEQ ID No. 29)
Anemonia sulcata	5'-CCTCTATCTTCATTCCTGC (SEQ ID No. 30)	5'-TATCTTCATTCCT GCGTAC (SEQ ID No. 31)
Discosoma sp. "magenta"	5'-TTCAGCACCCCATCACGAG (SEQ ID No. 32)	5'-ACGCTCAGAGCTG GGTTCC (SEQ ID No. 33)
Discosoma sp. "green"	5'-CCCTCAGCAATCCATCACGTT (SEQ ID No. 34)	5'-ATTATCTCAGTGGA TGGTTC (SEQ ID No. 35)

EXAMPLE 6**Expression of nFPs in *E.coli***

5 To prepare a DNA construct for novel fluorescent protein expression, two primers were synthesized for each cDNA: a 5'-directed "downstream" primer with the annealing site located in the 3'-UTR of the cDNA and a 3'-directed "upstream" primer corresponding to the site of translation start site (not including the first ATG codon) (Table
10 6). Both primers had 5'-heels coding for a site for a restriction endonuclease; in addition, the upstream primer was designed so as to allow the cloning of the PCR product into the pQE30 vector (Qiagen) in such a way that resulted in the fusion of reading frames of the vector-encoded 6xHis-tag and nFP. The PCR was performed as follows: 1 µl of
15 the 20-fold dilution of the amplified cDNA sample was added to a mixture containing 1x Advantage KlenTaq Polymerase Mix with buffer provided by the manufacturer (CLONTECH), 200 µM dNTPs, 0.2 µM of upstream primer and 0.2 µM of downstream primer, in a final total volume of 20 µl. The cycling profile was (Hybaid OmniGene
20 Thermocycler, tube control mode): 23-27 cycles for 95°C, 10 sec.; 60°C, 30 sec.; 72°C, 40 sec. The product of this amplification step was purified by phenol-chlorophorm extraction and ethanol precipitation and then cloned into pQE30 vector using restriction endonucleases corresponding to the primers' sequence according to standard
25 protocols.

All plasmids were amplified in XL-1 blue *E. coli* and purified by plasmid DNA miniprep kits (CLONTECH). The recombinant clones were selected by colony color, and grown in 3 ml of LB medium (supplemented with 100 µg/ml of ampicillin) at 37°C overnight. 100 µl

of the overnight culture was transferred into 200 ml of fresh LB medium containing 100 µg/ml of ampicillin and grown at 37°C, 200 rpm up to OD₆₀₀ 0.6-0.7. 1 mM IPTG was then added to the culture and incubation was allowed to proceed at 37°C for another 16 hours. The 5 cells were harvested and recombinant protein, which incorporated 6x His tags on the N-terminus, was purified using TALON™ metal-affinity resin according to the manufacturer's protocol (CLONTECH).

TABLE 6

Primers Used to Obtain Full Coding Region of nFPs for Cloning into Expression Construct

Species	Upstream Primer	Downstream Primer
Anemonia majano	5' -acatggatccgctttcaaaca agtttatc (SEQ ID No. 36) BamHI	5' -tagtactcgagcttattcgta ttcagtgaaatc (SEQ ID No. 37) XhoI
Clavularia sp.	L: 5'-acatggatccaacatttttga gaaacg (SEQ ID No. 38) BamHI S: 5'-acatggatccaaagctctaacc accatg (SEQ ID No. 39) BamHI	5' -tagtactcgagcaacacaa accctcagaccaa (SEQ ID No. 40) XhoI
Zoanthus sp.	5' - acatggatccgctcagtcaaag cacgg (SEQ ID No. 41) BamHI	5' -tagtactcgagggttggaaactacat tcttatca (SEQ ID No. 42) XhoI
Discosoma sp. "red"	5' - acatggatccaggccttccaagaat gttatac (SEQ ID No. 43) BamHI	5' -tagtactcgaggagccaagttc agcctta (SEQ ID No. 44) XhoI
Discosoma striata	5' - acatggatccaggcgttgtccaagagtg (SEQ ID No. 45) BamHI	5' -tagcgagcttatcatgcctc gtcacct (SEQ ID No. 46) SacI
Anemonia sulcata	5' - acatggatccgcttccttttaaagaagact (SEQ ID No. 47) BamHI	5' -tagtactcgagtcctggggagc ggcttg (SEQ ID No. 48) XhoI
Discosoma sp. "magenta"	5' - acatggatccaggcgttgtccaagaatgtgat (SEQ ID No. 49) BamHI	5' -tagtactcgaggccattacg ctaatac (SEQ ID No. 50) XhoI
Discosoma sp. "green"	5'-acatggatccaggcgtgcacttaaagaagaaatg (SEQ ID No. 51)	5'-tagtactcgagattcggttaat gccttg (SEQ ID No. 52)

EXAMPLE 7**Novel Fluorescent Proteins and cDNAs Encoding the Proteins**

Seven cDNA full-length cDNAs encoding fluorescent proteins were obtained (SEQ ID Nos. 45-51), and seven novel fluorescent proteins were produced (SEQ ID Nos. 53-59). The spectral properties of the isolated novel fluorescent proteins are shown in Table 7, and the emission and excitation spectra for the novel proteins are shown in Figures 3-11.

TABLE 7Spectral Properties of the Isolated NFPs.

Species	NFP Name	Abs. Max. n m	Emission Maximum n m	Maximum Extinction Coeff.	Relative Quantum Yield*	Relative Brightness **
Anemonia majano	amFP486	458	486	40,000	0.3	0.43
Clavularia sp.	cFP484	456	484	35,300	0.6	0.77
Zoanthus sp.	zFP506	496	506	35,600	0.79	1.02
Zoanthus sp.	zFP538	528	538	20,200	0.52	0.38
Discosoma sp. "red"	drFP583	558	583	22,500	0.29	0.24
Discosoma striata	dsFP483	443	483	23,900	0.57	0.50
Anemonia sulcata	asFP600	572	596	56,200	<0.001	-
Discosoma sp. "green"	dgFP512	502	512	20,360	0.3	0.21
Discosoma sp. "magenta"	dmFP592	573	593	21,800	0.11	0.09

5 *relative quantum yield was determined as compared to the quantum yield of *A. victoria* GFP.

**relative brightness is extinction coefficient multiplied by quantum yield divided by the same value for *A. victoria* GFP.

Multiple alignment of fluorescent proteins is shown in Figure 2A. The numbering is based on *Aequorea victoria* green fluorescent protein (GFP, SEQ ID No. 54). The amino acid sequences of the novel fluorescent proteins are labeled as SEQ ID Nos. 55-63. Two 5 proteins from *Zoanthus* and four from *Discosoma* are compared between each other: residues identical to the corresponding ones in the first protein of the series are represented by dashes. Introduced gaps are represented by dots. In the sequence of *A. victoria* GFP, the stretches forming β -sheets are underlined; the residues whose side 10 chains form the interior of the β -can are shaded. Figure 2B shows the N-terminal part of cFP484, which has no homology with the other proteins. The putative signal peptide is underlined.

The following references were cited herein.

1. Ormo et al., (1996) Science 273: 1392-1395.
- 15 2. Yang, F., et al., (1996) Nature Biotech 14: 1246-1251.
3. Cormack, et al., (1996) Gene 173, 33-38.
4. Haas, et al., (1996) Current Biology 6, 315-324.
5. Yang, et al., (1996) Nucleic Acids Research 24, 4592-4593.
6. Ghoda, et al.. (1990) J. Biol. Chem. 265: 11823-11826.
- 20 7. Prasher D.C. et al. (1992) Gene 111:229-33.
8. Kain et al. (1995) Biotechniques 19(4):650-55.
9. Chomczynski P., et al., (1987) Anal. Biochem. 162, 156-159.
10. Frohman et al., (1998) PNAS USA, 85, 8998-9002.

Any patents or publications mentioned in this specification 25 are indicative of the levels of those skilled in the art to which the invention pertains. These patents and publications are herein incorporated by reference to the same extent as if each individual publication was specifically and individually indicated to be incorporated by reference.

One skilled in the art will readily appreciate that the present invention is well adapted to carry out the objects and obtain the ends and advantages mentioned, as well as those inherent therein. The present examples along with the methods, procedures, treatments, 5 molecules, and specific compounds described herein are presently representative of preferred embodiments, are exemplary, and are not intended as limitations on the scope of the invention. Changes therein and other uses will occur to those skilled in the art which are encompassed within the spirit of the invention as defined by the scope 10 of the claims.

WHAT IS CLAIMED IS:

1. A method of identifying a DNA sequence encoding a fluorescent protein, comprising the step of:

5 screening for an existence of a nucleic acid sequence in a sample, wherein said nucleic acid sequence encodes a peptide having a sequence selected from the group consisting of SEQ ID Nos. 3, 5, 8, 11, 12 and 14, and wherein the existence of said nucleic acid sequence identifies the DNA sequence encoding the fluorescent protein.

10

2. A method of identifying a DNA sequence encoding a fluorescent protein, comprising the step of:

screening for an existence of a nucleic acid sequence in a sample, wherein said nucleic acid sequence hybridizes to a primer 15 selected from the group consisting of SEQ ID Nos. 4, 6, 7, 9, 10, 13, 15 and 16, and wherein the existence of said nucleic acid sequence identifies the DNA sequence encoding the fluorescent protein.

20

3. A method of analyzing a fluorescent protein in a cell,

comprising the steps of:

a) expressing a nucleic acid sequence encoding a fluorescent protein in said cell, wherein said protein having an amino acid sequence selected from the group consisting of SEQ ID Nos. 55-63; and

25

b) measuring a fluorescence signal from said protein.

4. The method of claim 3, further comprising the step of:

sorting said cell according to said signal.

5. The method of claim 4, wherein said step of sorting comprises sorting said cell by fluorescence activated cell sorting.

5 6. The method of claim 3, wherein said nucleic acid sequence comprises a gene of interest encoding a protein of interest fused to said fluorescent protein, wherein said protein of interest is distinct from said fluorescent protein.

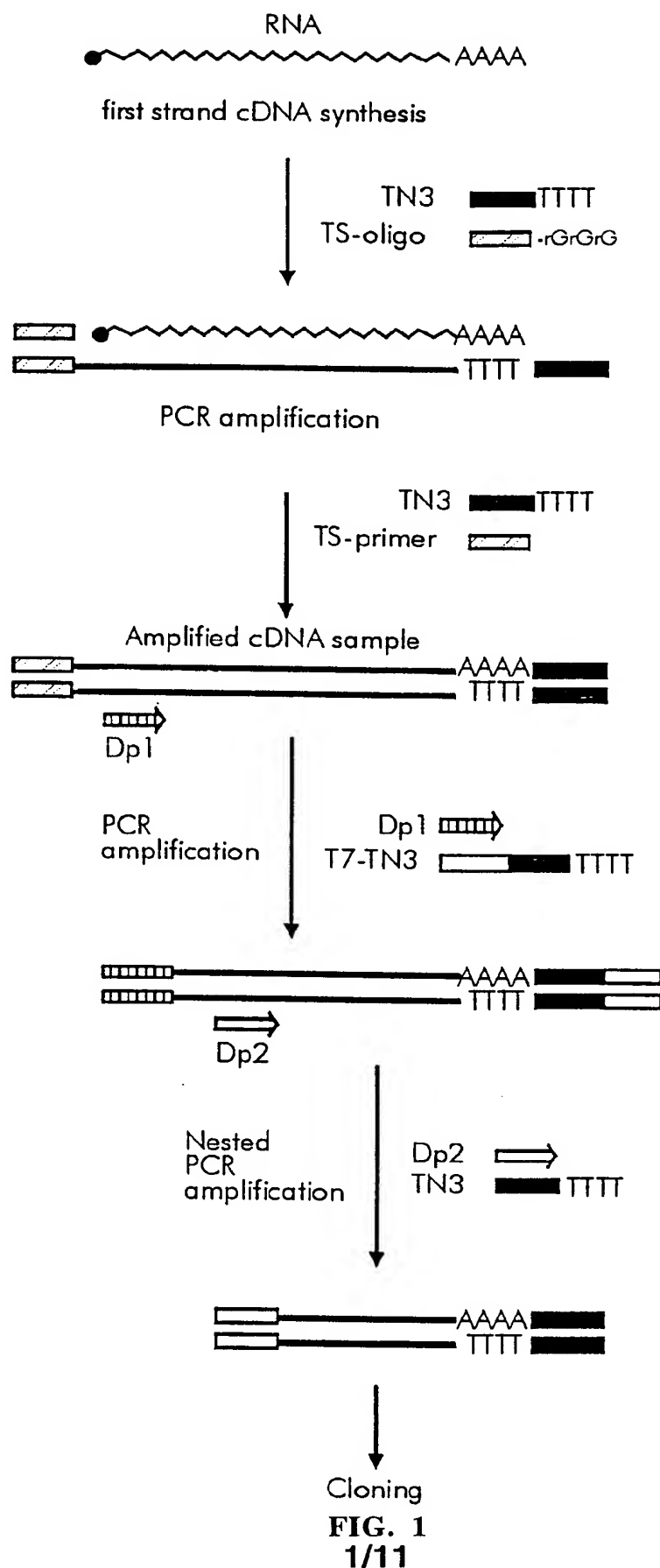
10 7. The method of claim 6, wherein the fluorescence signal indicates a presence of said gene of interest in said cell.

8. The method of claim 7, wherein said cell further comprises a protein of interest fused to said fluorescent protein.

15 9. The method of claim 8, further comprising the step of:

identifying an intracellular location of said fluorescent protein, thereby identifying an intracellular location of said protein of
20 interest.

10. An isolated and purified fluorescent protein selected from the group consisting of amFP486, cFP484, zFP506, zFP538, dsFP483, drFP583, asFP600, dgFP512 and dmFP592.



10	20	30	40	50		SEQ ID#
<u>MSKGEELFTG. VVPILVELGDVNNGHKFSVSGEGEGLDATYGKLTLKFIC</u> TT.GKLPVP..W GFP						54
MAQSKHGLTK. EMTMKYRMEGCVGDGHKFVITGEIGIGYPFKGKQAINLCVV..EGGPLPFAE					zFP506	57
--H----KE.-----H----N-----T-----I.-----S-					zFP538	58
MSWSKSVIKE. EMLIDLHLEGTFNNGHYFEIKGKGKGPNEGNTVTLEV..KGGPLPFGW					dsFP483	59
....M-AL--.Y-K-N-TM--VV--LP-K-R-D---YQ-SQEL--T-V.-----SY					dgFP512	62
-RS--N---.F-RFKVRM---V---E---E-E-R-Y---H---K-K-.----A-					drFP583	60
M-C--N---.F-RFKVRM---V---E---E-E-R-Y---HCS-K-M-.----AF					dmFP592	63
...MASFLKK.TMPFKTTIEGVNNGHYFKCTGKGEGNPFEGTQEMKIEVI..EGGPLPFAF					asFP600	61
MALSNKFIGD.DMKMTYIIMDCVNGHYFTVKGEGNKPYEGTQTSTFKVTMANGPLAFSF					amFP486	55
«KALTTMGVIKPDMKIKLKMEGNVNGHAFVIEGEGEKGPKYDGTHTLNLEVVKMAEGAPLPFSY					cFP484	56
60	70	80	90	100	110	
<u>PTLVTTFSYGVQCFSRYPDHMKQHDFFKSAM..:PEGYVQERTIFFKDDGNYKTRAEVKFEGD..</u>					GFP	
DILSAAFNYGNRVFTEYPQDIV..DYFKNSC...PAGYTWDRSFLFEDGAVCICNADITVSVEEN					zFP506	
----G-K--D-I-----G-----V-----K--					zFP538	
HILCPQFQYGNKAJVHHPPDDIP..DYLKLSF...PEGYTWERSMHFEDGGLCCINTNDISLTGN..					dsFP483	
D--TTM----R--NY-E---.IF-QTCSPGPN--S-Q-T-TY---V-TA-SN--VV-D..					dgFP512	
D--S----S-VY-K--A----K----.FK--V-N----VVTV-Q-S--QDG..					drFP583	
D--S----S-VY-K--A----K----.FK--V-N----VVTVSQ-S--KDG..					dmFP592	
HILSTSCMYGSKTFIKYVSGIP..DYFKQSF...PEGFTWERTTTYEDGGFLTAHQDTSLDGD..					asFP600	
DILSTVFKYGNRCFTAYPTSMP..DYFKQAF...PDGMSYERTFTYEDGGVATASWEISLKGN..					amFP486	
DILSNAFQYGNRALTKYPDDIA..DYFKQSF...PEGYSWERTMTFEDKGIVKVKSDISMEED..					cFP484	
120	130	140	150	160	170	
<u>TLVNRIELKGIDFKEDGNILGHKLEYNNYNSHNVI</u> MADKQKNGIKVNFKIRHNIEDGSVQL					GFP	
CMYHESKFYGVNFPAQGPVM.KKMTDNWEPSCEKIIIPVPKQGILKGDVSMYLLIKDGGRI.R					zFP506	
-I--K-I-N-M-----T--A----M-----Y-					zFP538	
CFNYDIKFTGLNFPPNGPVV.QKKTTGWEPESTERLYP..RDGVLIQDIHHALTVEGGGHYV					dsFP483	
T----H-M-A--LD--MM--R-MK----IMFE ---L-R-D-A-M-S-LLK----R					dgFP512	
--I-KV--I-V--SD--M---M--A----.K-E--K--KLKD---L					drFP583	
--I-EV--I-V--SD--M--RR-R----S----K---M--RL----L					dmFP592	
CLVYKVKILGNNFPADGPVM.QNKAGRWEPAEIVYE..VDGVLRGQSLMALCKPGGRHLT					asFP600	
CFEHKSTFHGVNFPAQGPVM.AKTTGWDPSEKMTV:.CDGILKGDVTAFLMLQGGGNR					amFP486	
SFIYEIRFDGMNFPNGPVM.QKKTLKWEPESTEIMYV,.RDGVLVGDISHSLLEGGGHYR					cFP484	
180	190	200	210	220	230	
<u>ADHYQONTPIGDG.PVLLPDNHYLSTOSALS</u> KDPNEKRDHMVLLEFVTAAGITHGMDELYK					GFP	
CQFDFTVYKAKSV..PRKMPDWHFIQHKLTREDRSDAKNQKWHLTEHAIASGSALP					zFP506	
-----S---E-----L-----Q-----FP---A					zFP538	
CDIKTVYDAKK...PVKMPGYHYVDTKLVIRSNDKEFM.KVEEHEIAVARHHPLQSQ					dsFP483	
--FE-I-KPN- V---D--F--HYIE-T-QQNYYN V--LT-V-E--YSS-EKIGKSKA					dgFP512	
VEF-SI-M----.QL--Y---S--D-T-HNEDYT.I--QY-RTEG--LFL					drFP583	
VEF-SI-MV-- PS-QL--Y---S--DMT-HNEDYT V--QY-KTQ----FIKPLQ					dmFP592	
CHLHTTYRSKKPASALKMPGFHFEDHRIEIMEEVEKGK.CYKQYEAAVGRYCDAAPSKLGHN					asFP600	
CQFHTSYKTKK...PVTMPPNVVHEHRIARTDLDKGGN.SVQLTEHAVAHITSVFPF					amFP486	
CDFKSIYKAKK...VVKLPDYHFVDHRIEILNHDKDYN.KVTLYENAVARYSLLPSQA					cFP484	

FIG. 2A

»

MKCKFVFCLSFLVLAITNANIFLRNEADLEEKTLRIP

FIG. 2B
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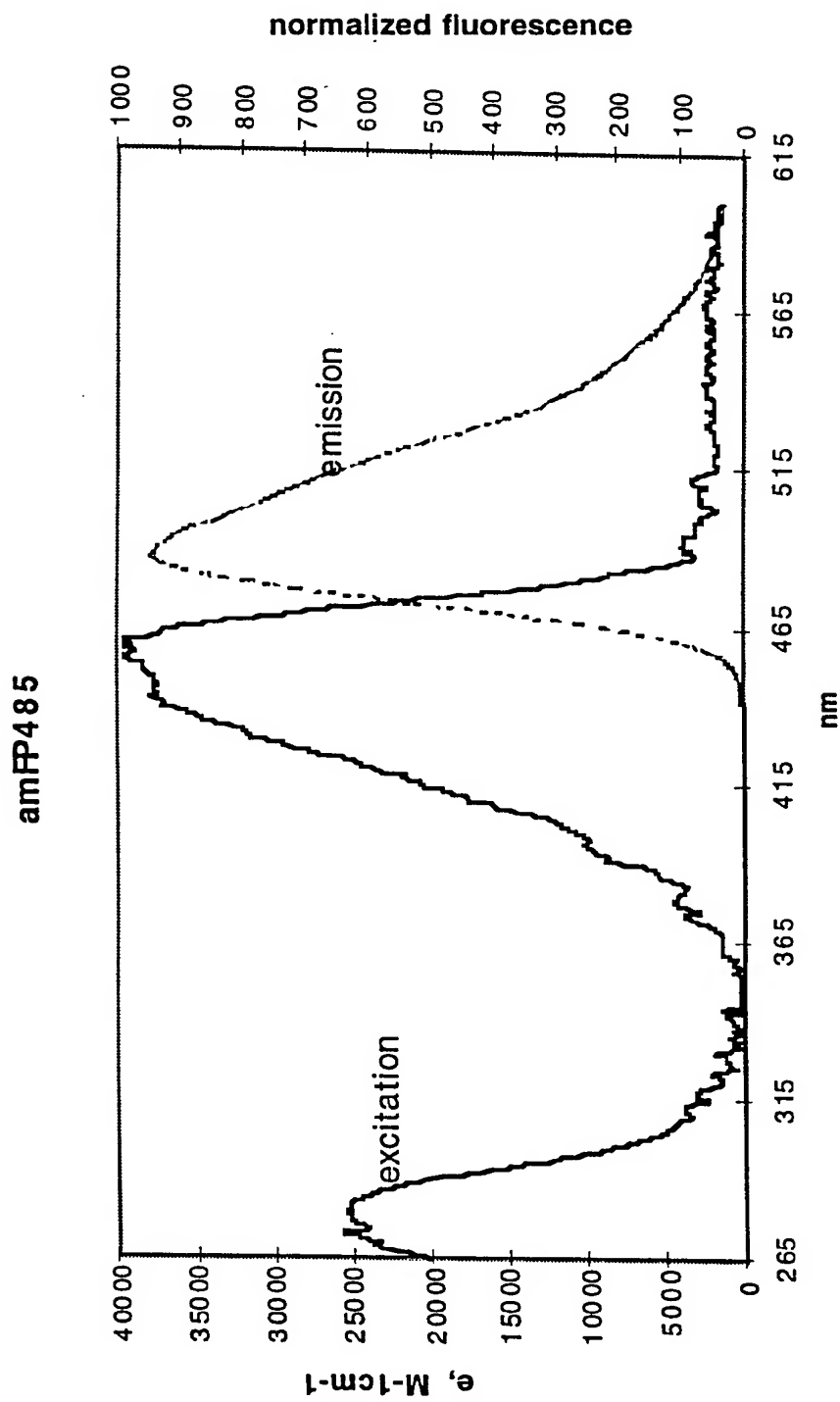


FIG. 3

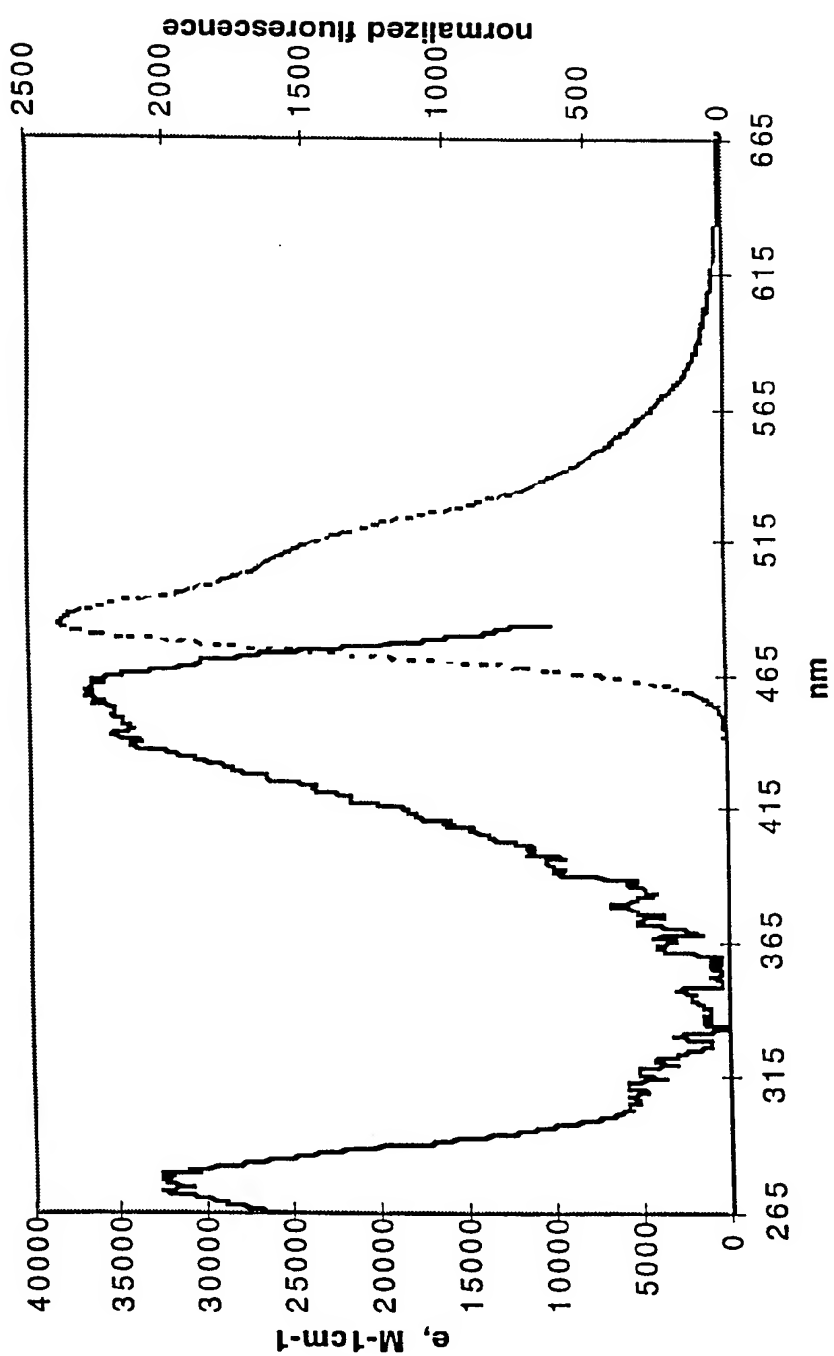
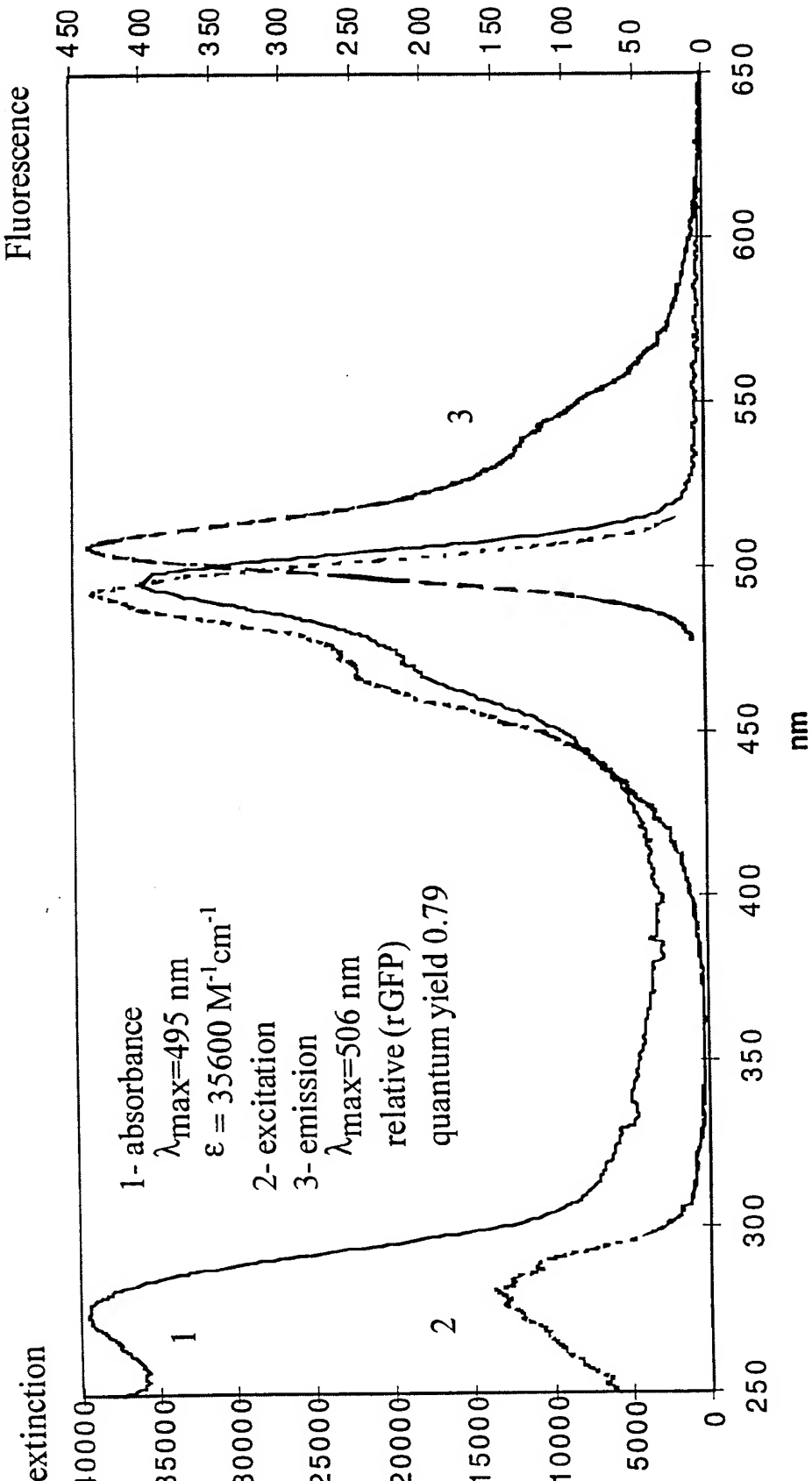
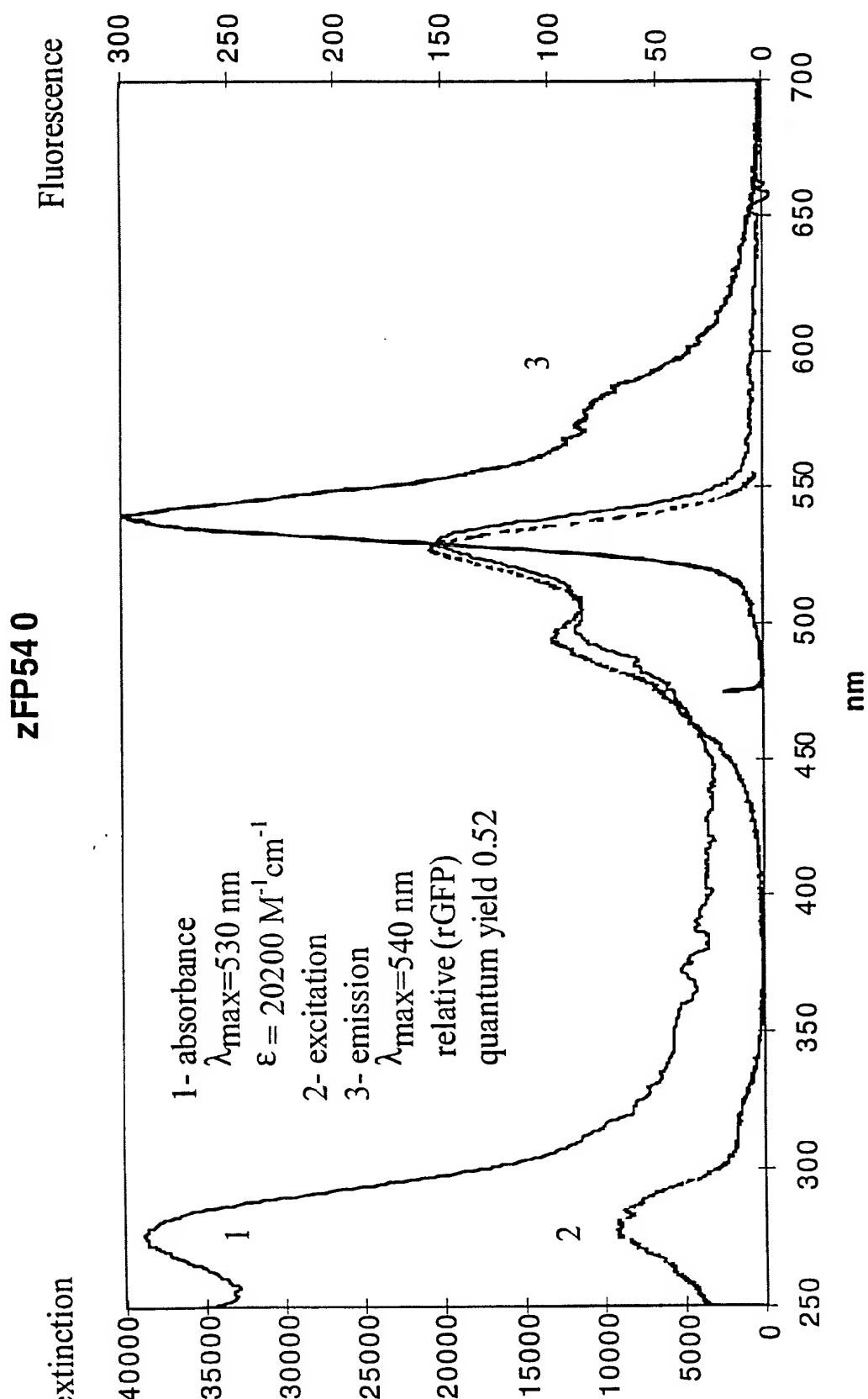


FIG. 4

zFP506

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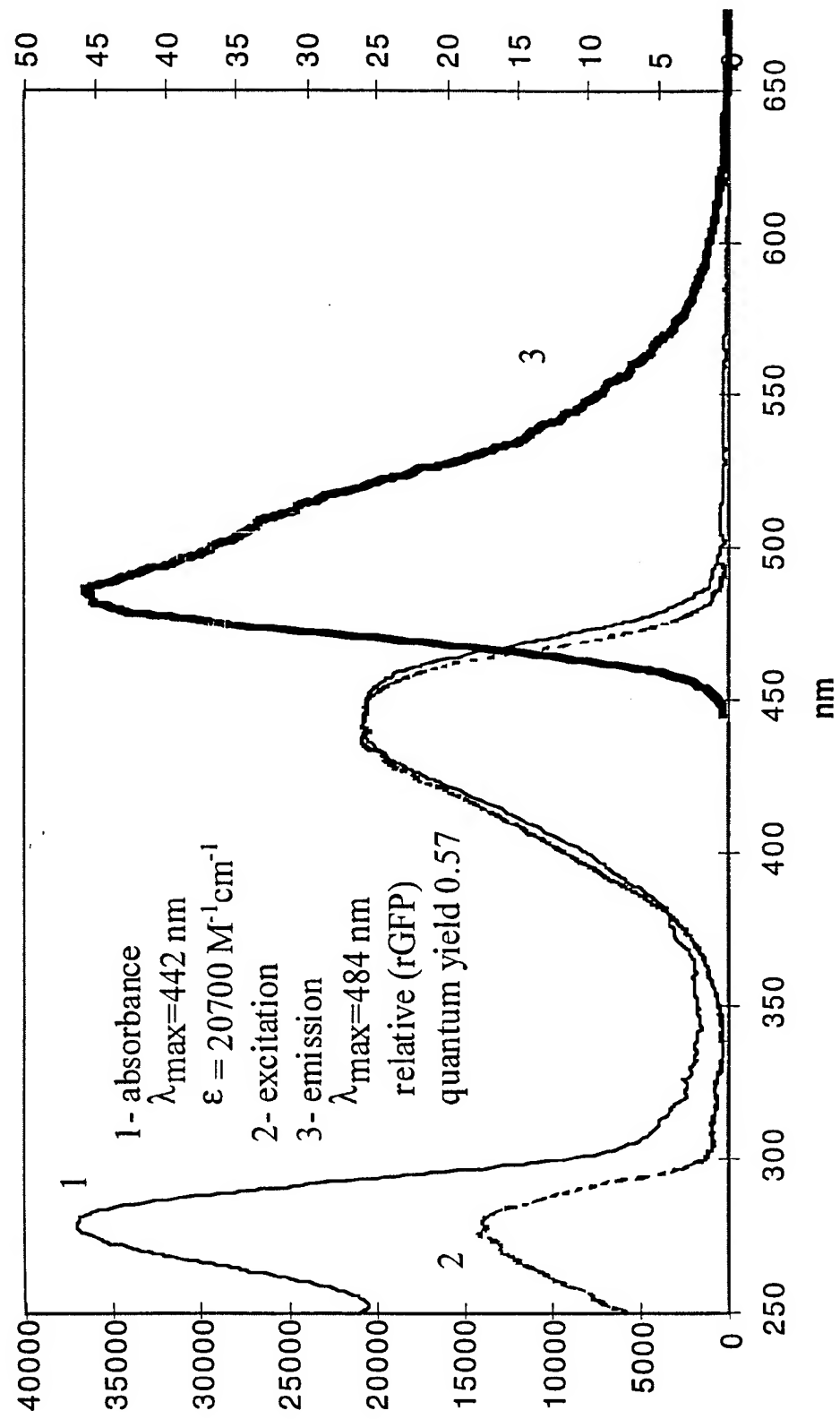
FIG. 5



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FIG. 6

dsFP484



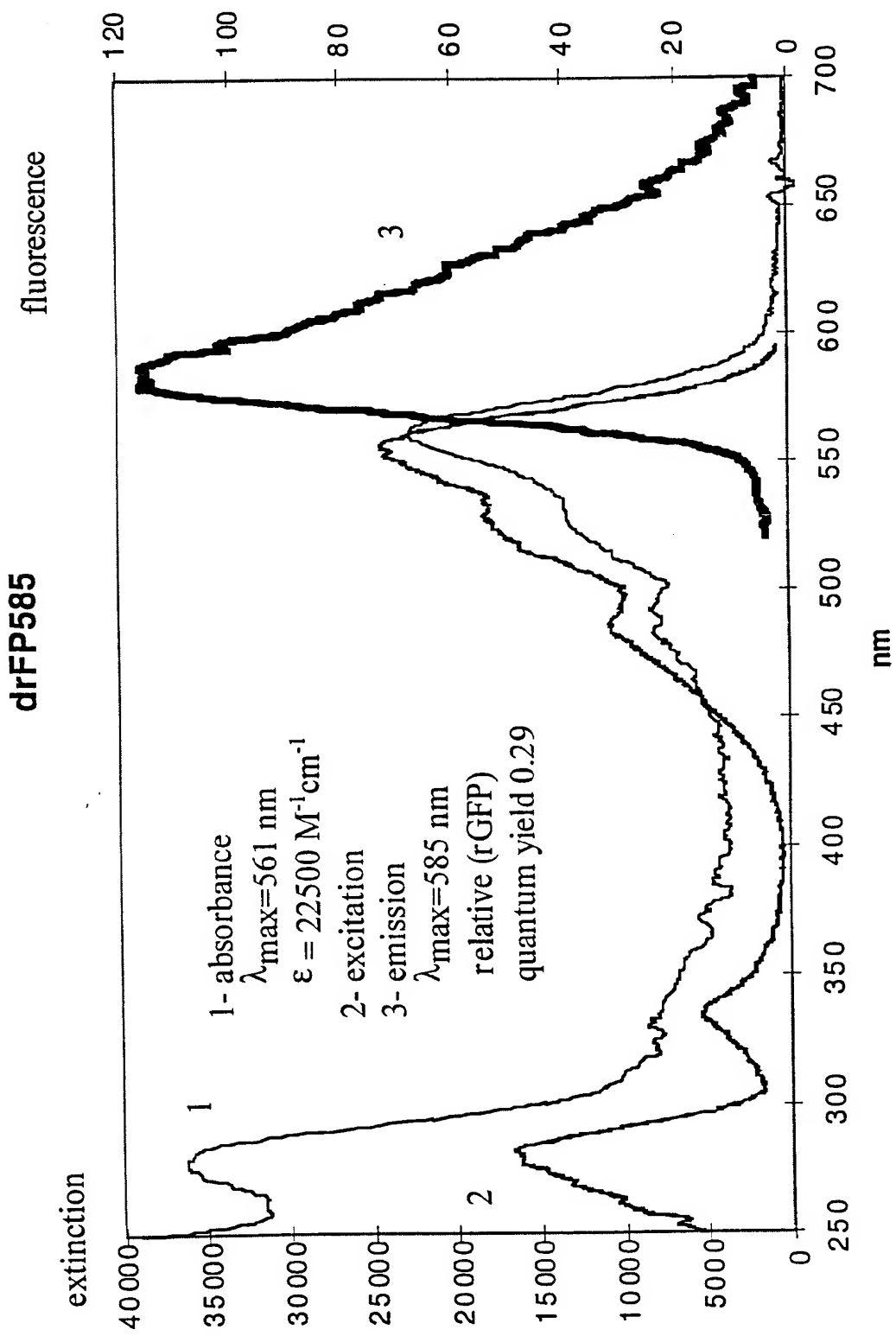
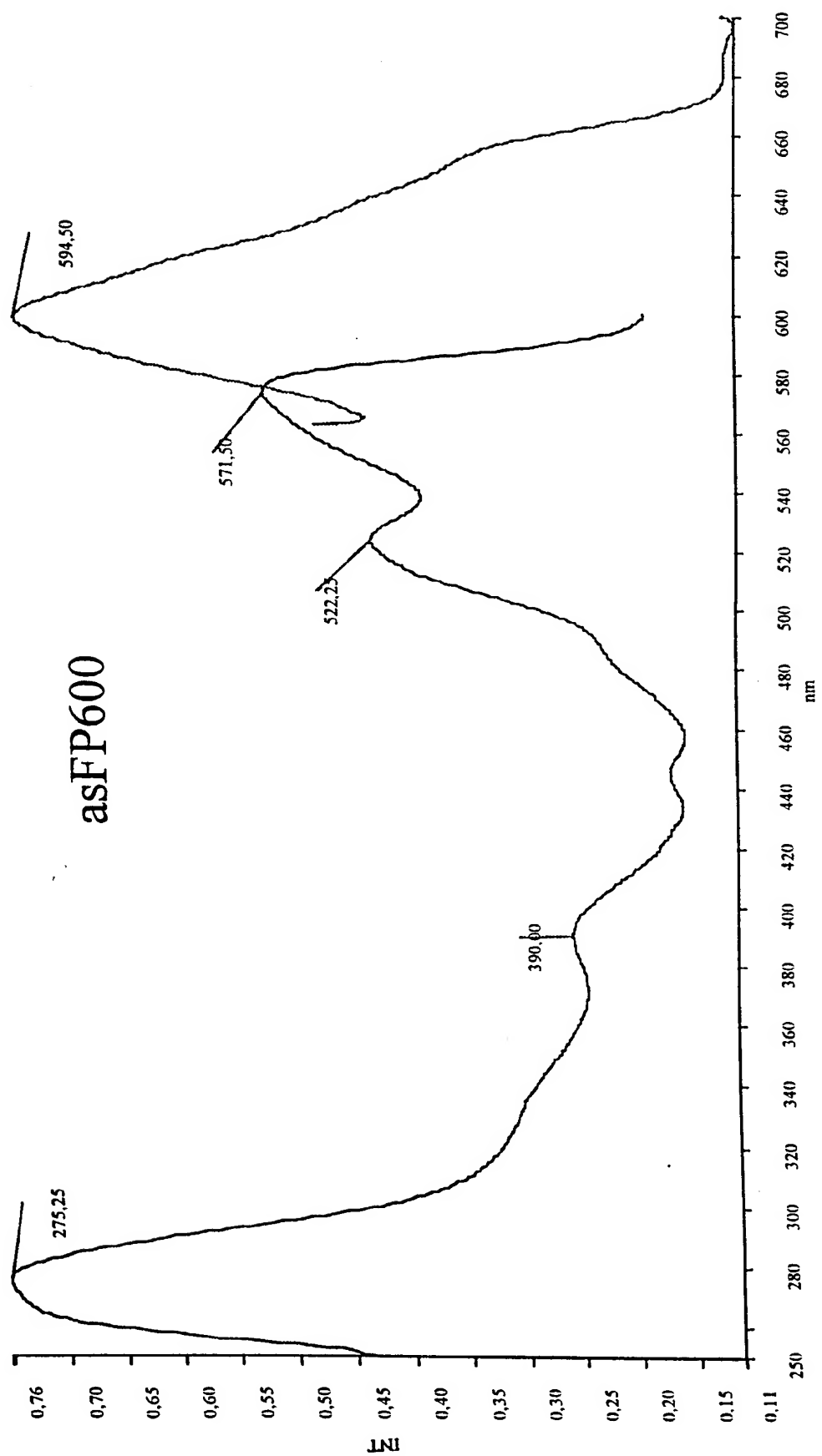


FIG. 8



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SUBSTITUTE SHEET (RULE 26)

FIG. 9

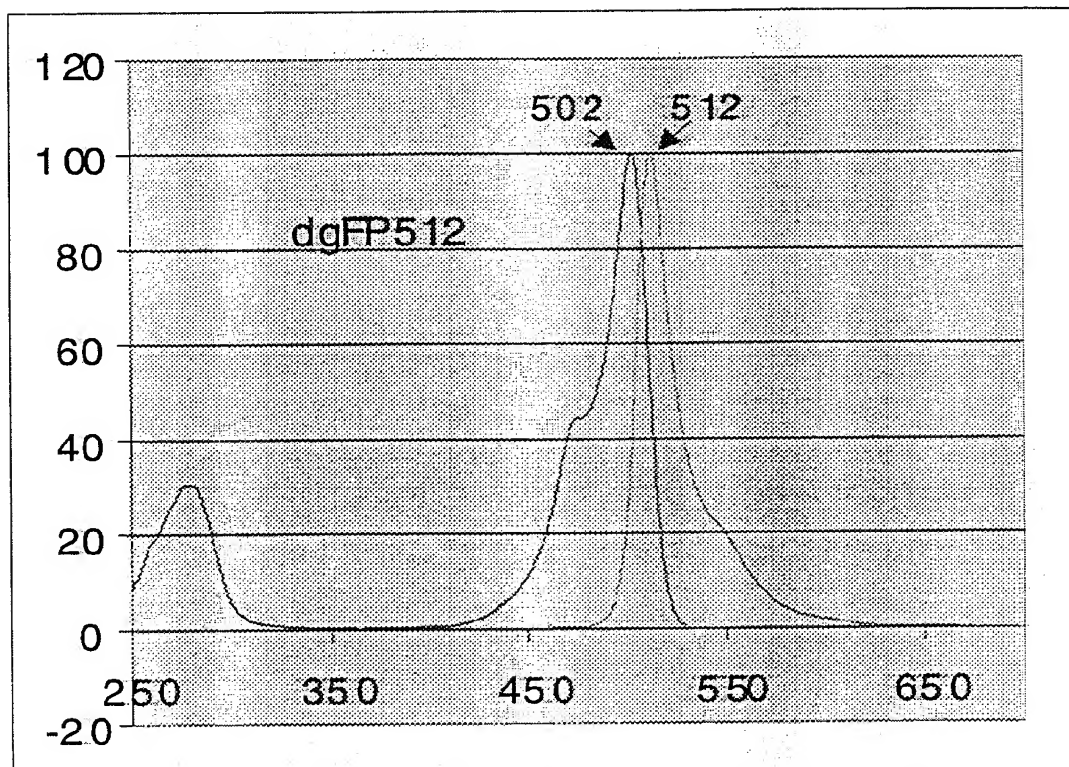


Fig. 10

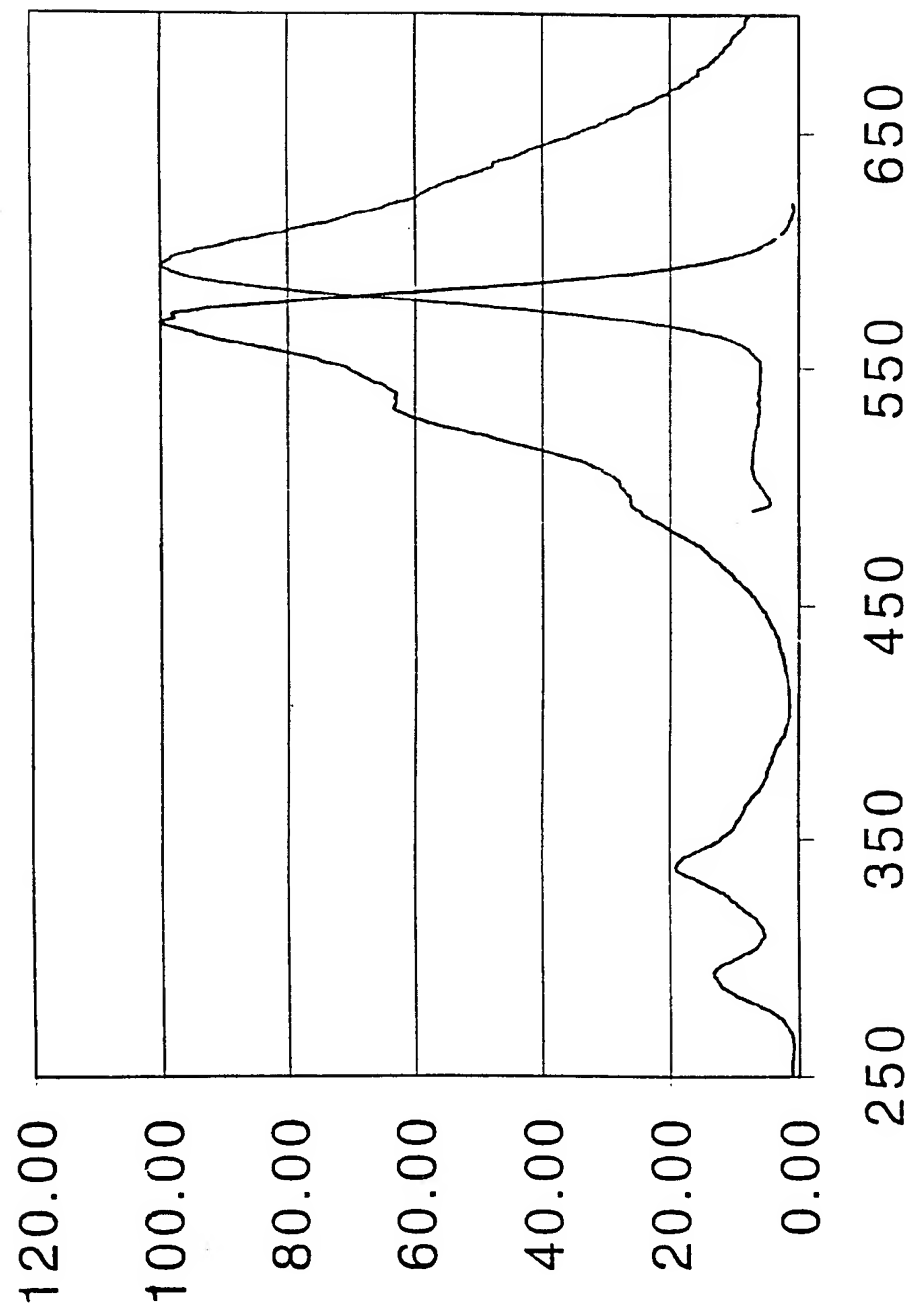


FIG. 11

SEQUENCE LISTING

<110> Lukyanov, Sergey A.
Labas, Yulii A.
Matz, Mikhail V.
5 Fradkov, Arcady F.

<120> Fluorescent proteins from non-bioluminescent
species of Class Anthozoa, genes encoding such
proteins and uses thereof

<130> D6196PCT

10 <141> 1999-12-10
<150> 09/210,330
<151> 1998-12-11
<160> 63

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35 <210> 3
<211> 6
<212> PRT

<213> *Aequorea victoria*
<220>
<222> 21
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represents unknown
<400> 3

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5

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15 <220>
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<223> primer NGH used for isolation of fluorescent
protein; n at position 12 represents any of the
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<400> 4

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20

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5

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protein
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<400> 7

gttacaggtg arggkgargg 20

20 <210> 8
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5

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5

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Gly Val Asn Phe Pro

5

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protein
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15

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20 <220>
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25 Gly Pro Val Met

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35 <223> primer PVMa used for isolation of fluorescent
protein; n at position 15 represents any of the

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<400>      15

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<211>      21
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<213>      artificial sequence
<220>
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<222>      15
<223>      primer PVMb used for isolation of fluorescent
            protein; n at position 15 represents any of the
            four bases
15      <400>      16

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20      <210>      17
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<220>
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30      <210>      18
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<213>      artificial sequence
<220>
<221>      primer_bind
<223>      primer T7-TS used in cDNA synthesis and RACE
35      <400>      18

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```

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<211> 22
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5 <213> artificial sequence
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<221> primer_bind
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<400> 19

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15 <213> artificial sequence
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20 <400> 20

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<210> 21
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30 *Anemonia majano*
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Zoanthus sp.

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<211> 19
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15 Discosoma sp. "red"
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Discosoma sp. "red"
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30 <210> 28
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Discosoma striata

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Discosoma sp. "magenta"
10 <400> 32

ttcagcaccc catcacgag 19

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20 <223> *Discosoma sp.* "magenta"
<400> 33

acgctcagag ctgggttcc 19

<210> 34
25 <211> 22
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Discosoma sp. "green"
<400> 34

ccctcagcaa tccatcacgt tc 22

35 <210> 35
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	<220>		
25	<221>	primer_bind	
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	tagtactcga gcttattcgt atttcagtga aatc		34
30	<210>	38	
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	<213>	artificial sequence	
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	<221>	primer_bind	
	<223>	upstream primer used to obtain full coding region	

of nFPs from *Clavularia* sp.

<400>	38	
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5 <210>	39	
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10 <221>	primer_bind	
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	of nFPs from <i>Clavularia</i> sp.	
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	region of nFPs from <i>Clavularia</i> sp.	
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25 tagtactcga gcaacacaaa ccctcagaca a		31
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	of nFPs from <i>Zoanthus</i> sp.	
35 <400>	41	
acatggatcc gtcagtc当地 agcacgg		28

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	<223>	downstream primer used to obtain full coding region of nFPs from <i>Zoanthus sp.</i>	
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	tagtactcgaa ggttgaaact acattcttat ca		32
	<210>	43	
	<211>	31	
15	<212>	DNA	
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	<221>	primer_bind	
	<223>	upstream primer used to obtain full coding region of nFPs from <i>Discosoma sp.</i> "red"	
20	<400>	43	
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	<210>	44	
25	<211>	29	
	<212>	DNA	
	<213>	artificial sequence	
	<220>		
	<221>	primer_bind	
30	<223>	downstream primer used to obtain full coding region of nFPs from <i>Discosoma sp.</i> "red"	
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	tagtactcgaa ggagccaaagt tcagcctta		29
35	<210>	45	
	<211>	28	
	<212>	DNA	

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	<220>		
	<221>	primer_bind	
5	<223>	upstream primer used to obtain full coding region of nFPs from <i>Discosoma striata</i>	
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10	<210>	46	
	<211>	28	
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	<220>		
	<221>	primer_bind	
15	<223>	downstream primer used to obtain full coding region of nFPs from <i>Discosoma striata</i>	
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	tagcgagctc tatcatgcct cgtcacct		28
20	<210>	47	
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25	<221>	primer_bind	
	<223>	upstream primer used to obtain full coding region of nFPs from <i>Anemonia sulcata</i>	
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	acatggatcc gtttccttt taaagaagac t		31
30	<210>	48	
	<211>	28	
	<212>	DNA	
	<213>	artificial sequence	
35	<220>		
	<221>	primer_bind	
	<223>	downstream primer used to obtain full coding	

region of nFPs from *Anemonia sulcata*

5	<400>	48	
	tagtactcg a gtccttggga gcggcttg		28
	<210>	49	
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	<220>		
10	<221>	primer_bind	
	<223>	upstream primer used to obtain full coding region of nFPs from <i>Discosoma sp.</i> "magenta"	
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	acatggatcc agttgttcca agaatgtgat		30
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	<210>	50	
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	<212>	DNA	
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20	<220>		
	<221>	primer_bind	
	<223>	downstream primer used to obtain full coding region of nFPs from <i>Discosoma sp.</i> "magenta"	
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25	tagtactcg a ggccattacg ctaatc		26
	<210>	51	
	<211>	31	
	<212>	DNA	
30	<213>	artificial sequence	
	<220>		
	<221>	primer_bind	
	<223>	upstream primer used to obtain full coding region of nFPs from <i>Discosoma sp.</i> "green"	
35	<400>	51	
	acatggatcc agtgcactta aagaagaaat g		31

<210> 52
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<212> DNA
5 <213> artificial sequence
<220>
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<223> downstream primer used to obtain full coding
region of nFPs from *Discosoma sp.* "green"
10 <400> 52

tagtactcga gattcggttt aatgccttg 29

<210> 53
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<223> TS-oligo used in cDNA synthesis and RACE
20 <400> 53

aagcagtggtaatcaacgcaggatcacgcrgrgrg 33

<210> 54
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25 <212> PRT
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30 Met Ser Lys Gly Glu Glu Leu Phe Thr Gly Val Val Pro Ile Leu
5 10 15
Val Glu Leu Asp Gly Asp Val Asn Gly His Lys Phe Ser Val Ser
20 25 30
Gly Glu Gly Glu Gly Asp Ala Thr Tyr Gly Lys Leu Thr Leu Lys
35 35 40 45
Phe Ile Cys Thr Thr Gly Lys Leu Pro Val Pro Trp Pro Thr Leu
50 55 60

	Val Thr Thr Phe Ser Tyr Gly Val Gln Cys Phe Ser Arg Tyr Pro		
	65	70	75
	Asp His Met Lys Gln His Asp Phe Phe Lys Ser Ala Met Pro Glu		
	80	85	90
5	Gly Tyr Val Gln Glu Arg Thr Ile Phe Phe Lys Asp Asp Gly Asn		
	95	100	105
	Tyr Lys Thr Arg Ala Glu Val Lys Phe Glu Gly Asp Thr Leu Val		
	110	115	120
	Asn Arg Ile Glu Leu Lys Gly Ile Asp Phe Lys Glu Asp Gly Asn		
10	125	130	135
	Ile Leu Gly His Lys Leu Glu Tyr Asn Tyr Asn Ser His Asn Val		
	140	145	150
	Tyr Ile Met Ala Asp Lys Gln Lys Asn Gly Ile Lys Val Asn Phe		
	155	160	165
15	Lys Ile Arg His Asn Ile Glu Asp Gly Ser Val Gln Leu Ala Asp		
	170	175	180
	His Tyr Gln Gln Asn Thr Pro Ile Gly Asp Gly Pro Val Leu Leu		
	185	190	195
	Pro Asp Asn His Tyr Leu Ser Thr Gln Ser Ala Leu Ser Lys Asp		
20	200	205	210
	Pro Asn Glu Lys Arg Asp His Met Val Leu Leu Glu Phe Val Thr		
	215	220	225
	Ala Ala Gly Ile Thr His Gly Met Asp Glu Leu Tyr Lys		
	230	235	
25	<210>	55	
	<211>	229	
	<212>	PRT	
	<213>	<i>Anemonia majano</i>	
30	<220>		
	<223>	amino acid sequence of amFP486	
	<400>	55	
	Met Ala Leu Ser Asn Lys Phe Ile Gly Asp Asp Met Lys Met Thr		
	5	10	15
35	Tyr His Met Asp Gly Cys Val Asn Gly His Tyr Phe Thr Val Lys		
	20	25	30
	Gly Glu Gly Asn Gly Lys Pro Tyr Glu Gly Thr Gln Thr Ser Thr		
	35	40	45

	Phe Lys Val Thr Met Ala Asn Gly Gly Pro Leu Ala Phe Ser Phe		
	50	55	60
	Asp Ile Leu Ser Thr Val Phe Lys Tyr Gly Asn Arg Cys Phe Thr		
	65	70	75
5	Ala Tyr Pro Thr Ser Met Pro Asp Tyr Phe Lys Gln Ala Phe Pro		
	80	85	90
	Asp Gly Met Ser Tyr Glu Arg Thr Phe Thr Tyr Glu Asp Gly Gly		
	95	100	105
	Val Ala Thr Ala Ser Trp Glu Ile Ser Leu Lys Gly Asn Cys Phe		
10	110	115	120
	Glu His Lys Ser Thr Phe His Gly Val Asn Phe Pro Ala Asp Gly		
	125	130	135
	Pro Val Met Ala Lys Lys Thr Thr Gly Trp Asp Pro Ser Phe Glu		
	140	145	150
15	Lys Met Thr Val Cys Asp Gly Ile Leu Lys Gly Asp Val Thr Ala		
	155	160	165
	Phe Leu Met Leu Gln Gly Gly Asn Tyr Arg Cys Gln Phe His		
	170	175	180
	Thr Ser Tyr Lys Thr Lys Pro Val Thr Met Pro Pro Asn His		
20	185	190	195
	Val Val Glu His Arg Ile Ala Arg Thr Asp Leu Asp Lys Gly Gly		
	200	205	210
	Asn Ser Val Gln Leu Thr Glu His Ala Val Ala His Ile Thr Ser		
	215	220	225
25	Val Val Pro Phe		

	<210>	56	
	<211>	266	
30	<212>	PRT	
	<213>	<i>Clavularia sp.</i>	
	<220>		
	<223>	amino acid sequence of cFP484	
	<400>	56	
35	Met Lys Cys Lys Phe Val Phe Cys Leu Ser Phe Leu Val Leu Ala		
	5	10	15
	Ile Thr Asn Ala Asn Ile Phe Leu Arg Asn Glu Ala Asp Phe Glu		

	20	25	30
10	Glu Lys Thr Phe Arg Ile Pro Lys Ala Leu Thr Thr Met Gly Val		
	35	40	45
5	Ile Lys Pro Asp Met Lys Ile Lys Leu Lys Met Glu Gly Asn Val		
	50	55	60
	Asn Gly His Ala Phe Val Ile Glu Gly Glu Gly Glu Lys Pro		
	65	70	75
	Tyr Asp Gly Thr His Thr Leu Asn Leu Glu Val Lys Glu Gly Ala		
	80	85	90
10	Pro Leu Pro Phe Ser Tyr Asp Ile Leu Ser Asn Ala Phe Gln Tyr		
	95	100	105
	Gly Asn Arg Ala Leu Thr Lys Tyr Pro Asp Asp Ile Ala Asp Tyr		
	110	115	120
15	Phe Lys Gln Ser Phe Pro Glu Gly Tyr Ser Trp Glu Arg Thr Met		
	125	130	135
	Thr Phe Glu Asp Lys Gly Ile Val Lys Val Lys Ser Asp Ile Ser		
	140	145	150
	Met Glu Glu Asp Ser Phe Ile Tyr Glu Ile Arg Phe Asp Gly Met		
	155	160	165
20	Asp Phe Pro Pro Asn Gly Pro Val Met Gln Lys Lys Thr Leu Lys		
	170	175	180
	Trp Glu Pro Ser Thr Glu Ile Met Tyr Val Arg Asp Gly Val Leu		
	185	190	195
	Val Gly Asp Ile Ser His Ser Leu Leu Leu Glu Gly Gly His		
25	200	205	210
	Tyr Arg Cys Asp Phe Lys Ser Ile Tyr Lys Ala Lys Lys Val Val		
	215	220	225
	Lys Leu Pro Asp Tyr His Phe Val Asp His Arg Ile Glu Ile Leu		
	230	235	240
30	Asn His Asp Lys Asp Tyr Asn Lys Val Thr Leu Tyr Glu Asn Ala		
	245	250	255
	Val Ala Arg Tyr Ser Leu Leu Pro Ser Gln Ala		
	260	265	
35	<210>	57	
	<211>	230	
	<212>	PRT	

<213> *Zoanthus sp.*
 <220>
 <223> amino acid sequence of zFP506
 <400> 57

5	Ala Gln Ser Lys His Gly Leu Thr Lys Glu Met Thr Met Lys Tyr		
	5	10	15
	Arg Met Glu Gly Cys Val Asp Gly His Lys Phe Val Ile Thr Gly		
	20	25	30
	Glu Gly Ile Gly Tyr Pro Phe Lys Gly Lys Gln Ala Ile Asn Leu		
10	35	40	45
	Cys Val Val Glu Gly Pro Leu Pro Phe Ala Glu Asp Ile Leu		
	50	55	60
	Ser Ala Ala Phe Asn Tyr Gly Asn Arg Val Phe Thr Glu Tyr Pro		
	65	70	75
15	Gln Asp Ile Val Asp Tyr Phe Lys Asn Ser Cys Pro Ala Gly Tyr		
	80	85	90
	Thr Trp Asp Arg Ser Phe Leu Phe Glu Asp Gly Ala Val Cys Ile		
	95	100	105
	Cys Asn Ala Asp Ile Thr Val Ser Val Glu Glu Asn Cys Met Tyr		
20	110	115	120
	His Glu Ser Lys Phe Tyr Gly Val Asn Phe Pro Ala Asp Gly Pro		
	125	130	135
	Val Met Lys Lys Met Thr Asp Asn Trp Glu Pro Ser Cys Glu Lys		
	140	145	150
25	Ile Ile Pro Val Pro Lys Gln Gly Ile Leu Lys Gly Asp Val Ser		
	155	160	165
	Met Tyr Leu Leu Lys Asp Gly Gly Arg Leu Arg Cys Gln Phe		
	170	175	180
	Asp Thr Val Tyr Lys Ala Lys Ser Val Pro Arg Lys Met Pro Asp		
30	185	190	195
	Trp His Phe Ile Gln His Lys Leu Thr Arg Glu Asp Arg Ser Asp		
	200	205	210
	Ala Lys Asn Gln Lys Trp His Leu Thr Glu His Ala Ile Ala Ser		
	215	220	225
35	Gly Ser Ala Leu Pro		
	230		

	<210>	58	
	<211>	230	
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5	<213>	Zoanthus sp.	
	<220>		
	<223>	amino acid sequence of zFP538	
	<400>	58	
	Met Ala His Ser Lys His Gly Leu Lys Glu Glu Met Thr Met Lys		
10	5	10	15
	Tyr His Met Glu Gly Cys Val Asn Gly His Lys Phe Val Ile Thr		
	20	25	30
	Gly Glu Gly Ile Gly Tyr Pro Phe Lys Gly Lys Gln Thr Ile Asn		
	35	40	45
15	Leu Cys Val Ile Glu Gly Gly Pro Leu Pro Phe Ser Glu Asp Ile		
	50	55	60
	Leu Ser Ala Gly Phe Lys Tyr Gly Asp Arg Ile Phe Thr Glu Tyr		
	65	70	75
	Pro Gln Asp Ile Val Asp Tyr Phe Lys Asn Ser Cys Pro Ala Gly		
20	80	85	90
	Tyr Thr Trp Gly Ser Phe Leu Phe Glu Asp Gly Ala Val Cys Ile		
	95	100	105
	Cys Asn Val Asp Ile Thr Val Ser Val Lys Glu Asn Cys Ile Tyr		
	110	115	120
25	His Lys Ser Ile Phe Asn Gly Met Asn Phe Pro Ala Asp Gly Pro		
	125	130	135
	Val Met Lys Lys Met Thr Thr Asn Trp Glu Ala Ser Cys Glu Lys		
	140	145	150
	Ile Met Pro Val Pro Lys Gln Gly Ile Leu Lys Gly Asp Val Ser		
30	155	160	165
	Met Tyr Leu Leu Leu Lys Asp Gly Gly Arg Tyr Arg Cys Gln Phe		
	170	175	180
	Asp Thr Val Tyr Lys Ala Lys Ser Val Pro Ser Lys Met Pro Glu		
	185	190	195
35	Trp His Phe Ile Gln His Lys Leu Leu Arg Glu Asp Arg Ser Asp		
	200	205	210
	Ala Lys Asn Gln Lys Trp Gln Leu Thr Glu His Ala Ile Ala Phe		
	215	220	225

Pro Ser Ala Leu Ala

230

5	<210>	59
	<211>	232
	<212>	PRT
	<213>	<i>Discosoma striata</i>
	<220>	
10	<223>	amino acid sequence of dsFP483
	<400>	59
	Met Ser Cys Ser Lys Ser Val Ile Lys Glu Glu Met Leu Ile Asp	
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	Leu His Leu Glu Gly Thr Phe Asn Gly His Tyr Phe Glu Ile Lys	
15	20	25 30
	Gly Lys Gly Lys Gly Gln Pro Asn Glu Gly Thr Asn Thr Val Thr	
	35	40 45
	Leu Glu Val Thr Lys Gly Gly Pro Leu Pro Phe Gly Trp His Ile	
	50	55 60
20	Leu Cys Pro Gln Phe Gln Tyr Gly Asn Lys Ala Phe Val His His	
	65	70 75
	Pro Asp Asn Ile His Asp Tyr Leu Lys Leu Ser Phe Pro Glu Gly	
	80	85 90
	Tyr Thr Trp Glu Arg Ser Met His Phe Glu Asp Gly Gly Leu Cys	
25	95	100 105
	Cys Ile Thr Asn Asp Ile Ser Leu Thr Gly Asn Cys Phe Tyr Tyr	
	110	115 120
	Asp Ile Lys Phe Thr Gly Leu Asn Phe Pro Pro Asn Gly Pro Val	
	125	130 135
30	Val Gln Lys Lys Thr Thr Gly Trp Glu Pro Ser Thr Glu Arg Leu	
	140	145 150
	Tyr Pro Arg Asp Gly Val Leu Ile Gly Asp Ile His His Ala Leu	
	155	160 165
	Thr Val Glu Gly Gly His Tyr Ala Cys Asp Ile Lys Thr Val	
35	170	175 180
	Tyr Arg Ala Lys Lys Ala Ala Leu Lys Met Pro Gly Tyr His Tyr	
	185	190 195

	Val Asp Thr Lys Leu Val Ile Trp Asn Asn Asp Lys Glu Phe Met		
	200	205	210
	Lys Val Glu Glu His Glu Ile Ala Val Ala Arg His His Pro Phe		
	215	220	225
5	Tyr Glu Pro Lys Lys Asp Lys		
	230		
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10	<212>	PRT	
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	<220>		
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	<400>	60	
15	Met Arg Ser Ser Lys Asn Val Ile Lys Glu Phe Met Arg Phe Lys		
	5	10	15
	Val Arg Met Glu Gly Thr Val Asn Gly His Glu Phe Glu Ile Glu		
	20	25	30
	Gly Glu Gly Glu Gly Arg Pro Tyr Glu Gly His Asn Thr Val Lys		
20	35	40	45
	Leu Lys Val Thr Lys Gly Gly Pro Leu Pro Phe Ala Trp Asp Ile		
	50	55	60
	Leu Ser Pro Gln Phe Gln Tyr Gly Ser Lys Val Tyr Val Lys His		
	65	70	75
25	Pro Ala Asp Ile Pro Asp Tyr Lys Lys Leu Ser Phe Pro Glu Gly		
	80	85	90
	Phe Lys Trp Glu Arg Val Met Asn Phe Glu Asp Gly Gly Val Val		
	95	100	105
	Thr Val Thr Gln Asp Ser Ser Leu Gln Asp Gly Cys Phe Ile Tyr		
30	110	115	120
	Lys Val Lys Phe Ile Gly Val Asn Phe Pro Ser Asp Gly Pro Val		
	125	130	135
	Met Gln Lys Lys Thr Met Gly Trp Glu Ala Ser Thr Glu Arg Leu		
	140	145	150
35	Tyr Pro Arg Asp Gly Val Leu Lys Gly Glu Ile His Lys Ala Leu		
	155	160	165
	Lys Leu Lys Asp Gly Gly His Tyr Leu Val Glu Phe Lys Ser Ile		
	170	175	180

	Tyr Met Ala Lys Lys Pro Val Gln Leu Pro Gly Tyr Tyr Tyr Val		
	185	190	195
	Asp Ser Lys Leu Asp Ile Thr Ser His Asn Glu Asp Tyr Thr Ile		
	200	205	210
5	Val Glu Gln Tyr Glu Arg Thr Glu Gly Arg His His Leu Phe Leu		
	215	220	225
	<210> 61		
	<211> 232		
10	<212> PRT		
	<213> <i>Anemonia sulcata</i>		
	<220>		
	<223> amino acid sequence of asFP600		
	<400> 61		
15	Met Ala Ser Phe Leu Lys Lys Thr Met Pro Phe Lys Thr Thr Ile		
	5	10	15
	Glu Gly Thr Val Asn Gly His Tyr Phe Lys Cys Thr Gly Lys Gly		
	20	25	30
	Glu Gly Asn Pro Phe Glu Gly Thr Gln Glu Met Lys Ile Glu Val		
20	35	40	45
	Ile Glu Gly Gly Pro Leu Pro Phe Ala Phe His Ile Leu Ser Thr		
	50	55	60
	Ser Cys Met Tyr Gly Ser Lys Thr Phe Ile Lys Tyr Val Ser Gly		
	65	70	75
25	Ile Pro Asp Tyr Phe Lys Gln Ser Phe Pro Glu Gly Phe Thr Trp		
	80	85	90
	Glu Arg Thr Thr Tyr Glu Asp Gly Gly Phe Leu Thr Ala His		
	95	100	105
	Gln Asp Thr Ser Leu Asp Gly Asp Cys Leu Val Tyr Lys Val Lys		
30	110	115	120
	Ile Leu Gly Asn Asn Phe Pro Ala Asp Gly Pro Val Met Gln Asn		
	125	130	135
	Lys Ala Gly Arg Trp Glu Pro Ala Thr Glu Ile Val Tyr Glu Val		
	140	145	150
35	Asp Gly Val Leu Arg Gly Gln Ser Leu Met Ala Leu Lys Cys Pro		
	155	160	165
	Gly Gly Arg His Leu Thr Cys His Leu His Thr Thr Tyr Arg Ser		
	170	175	180

	Lys Lys Pro Ala Ser Ala Leu Lys Met Pro Gly Phe His Phe Glu		
	185	190	195
	Asp His Arg Ile Glu Ile Met Glu Glu Val Glu Lys Gly Lys Cys		
	200	205	210
5	Tyr Lys Gln Tyr Glu Ala Ala Val Gly Arg Tyr Cys Asp Ala Ala		
	215	220	225
	Pro Ser Lys Leu Gly His Asn		
	230		
10	<210>	62	
	<211>	231	
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	<213>	<i>Discosoma sp. "green"</i>	
	<220>		
15	<223>	amino acid sequence of dgFP512	
	<400>	62	
	Met Ser Ala Leu Lys Glu Glu Met Lys Ile Asn Leu Thr Met Glu		
	5	10	15
	Gly Val Val Asn Gly Leu Pro Phe Lys Ile Arg Gly Asp Gly Lys		
20	20	25	30
	Gly Lys Pro Tyr Gln Gly Ser Gln Glu Leu Thr Leu Thr Val Val		
	35	40	45
	Lys Gly Gly Pro Leu Pro Phe Ser Tyr Asp Ile Leu Thr Thr Met		
	50	55	60
25	Phe Gln Tyr Gly Asn Arg Ala Phe Val Asn Tyr Pro Glu Asp Ile		
	65	70	75
	Pro Asp Ile Phe Lys Gln Thr Cys Ser Gly Pro Asn Gly Gly Tyr		
	80	85	90
	Ser Trp Gln Arg Thr Met Thr Tyr Glu Asp Gly Gly Val Cys Thr		
30	95	100	105
	Ala Thr Ser Asn Ile Ser Val Val Gly Asp Thr Phe Asn Tyr Asp		
	110	115	120

	Ile His Phe Met Gly Ala Asn Phe Pro Leu Asp Gly Pro Val Met		
	125	130	135
	Gln Lys Arg Thr Met Lys Trp Glu Pro Ser Thr Glu Ile Met Phe		
	140	145	150
5	Glu Arg Asp Gly Met Leu Arg Gly Asp Ile Ala Met Ser Leu Leu		
	155	160	165
	Leu Lys Gly Gly His Tyr Arg Cys Asp Phe Glu Thr Ile Tyr		
	170	175	180
	Lys Pro Asn Lys Val Val Lys Met Pro Asp Tyr His Phe Val Asp		
10	185	190	195
	His Cys Ile Glu Ile Thr Ser Gln Gln Asp Tyr Tyr Asn Val Val		
	200	205	210
	Glu Leu Thr Glu Val Ala Glu Ala Arg Tyr Ser Ser Leu Glu Lys		
	215	220	225
15	Ile Gly Lys Ser Lys Ala		
	230		
	<210> 63		
	<211> 235		
20	<212> PRT		
	<213> <i>Discosoma sp. "magenta"</i>		
	<220>		
	<223> amino acid sequence of dmFP592		
	<400> 63		
25	Met Ser Cys Ser Lys Asn Val Ile Lys Glu Phe Met Arg Phe Lys		
	5	10	15
	Val Arg Met Glu Gly Thr Val Asn Gly His Glu Phe Glu Ile Lys		
	20	25	30
	Gly Glu Gly Glu Gly Arg Pro Tyr Glu Gly His Cys Ser Val Lys		
30	35	40	45

Leu Met Val Thr Lys Gly Gly Pro Leu Pro Phe Ala Phe Asp Ile
50 55 60

Leu Ser Pro Gln Phe Gln Tyr Gly Ser Lys Val Tyr Val Lys His
65 70 75

5 Pro Ala Asp Ile Pro Asp Tyr Lys Lys Leu Ser Phe Pro Glu Gly
80 85 90

Phe Lys Trp Glu Arg Val Met Asn Phe Glu Asp Gly Gly Val Val
100 105 110

Thr Val Ser Gln Asp Ser Ser Leu Lys Asp Gly Cys Phe Ile Tyr
10 115 120 125

Glu Val Lys Phe Ile Gly Val Asn Phe Pro Ser Asp Gly Pro Val
130 135 140

Met Gln Arg Arg Thr Arg Gly Trp Glu Ala Ser Ser Glu Arg Leu
145 150 155

15 Tyr Pro Arg Asp Gly Val Leu Lys Gly Asp Ile His Met Ala Leu
160 165 170

Arg Leu Glu Gly Gly His Tyr Leu Val Glu Phe Lys Ser Ile
175 180 185

Tyr Met Val Lys Lys Pro Ser Val Gln Leu Pro Gly Tyr Tyr Tyr
20 190 195 200

Val Asp Ser Lys Leu Asp Met Thr Ser His Asn Glu Asp Tyr Thr
205 210 215

Val Val Glu Gln Tyr Glu Lys Thr Gln Gly Arg His His Pro Phe
220 225 230

25 Ile Lys Pro Leu Gln
235

INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/29405

A. CLASSIFICATION OF SUBJECT MATTER

IPC(7) :C12Q 1/68; C07K 14/435
US CL :435/6, 69.1; 530/350

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/6, 69.1, 968; 530/350; 424/9.6, 436/172

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

Please See Extra Sheet.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
***	The sequence diskette submitted with the description was defective; thus the references listed below were obtained solely by a WORD search, and not by a search of the SEQ ID NOs.	***
X, P	MATZ et al. Fluorescent proteins from nonbioluminescent Anthozoa species. Nature Biotechnology. October 1999, Volume 17, No. 10, pages 969-973, entire document.	1-10
X, P	DE 197 18 640 A1 (WIEDENMANN) 22 July 1999, entire document.	3-10

Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents:	"T"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"A" document defining the general state of the art which is not considered to be of particular relevance		
"E" earlier document published on or after the international filing date	"X"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"Y"	document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"O" document referring to an oral disclosure, use, exhibition or other means	"&"	document member of the same patent family
"P" document published prior to the international filing date but later than the priority date claimed		

Date of the actual completion of the international search
18 FEBRUARY 2000

Date of mailing of the international search report

02 MAR 2000

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